U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE FORM PTO-1390 (REV 12-99) ST98009 TRANSMITTAL LETTER TO THE UNITED STATES U.S. APPLICATION NO DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371 INTERNATIONAL APPLICATION NO. INTERNATIONAL FILING DATE PRIORITY DATE CI PCT/FR99/00740 -30 March 1999 -02 April 1998 TITLE OF INVENTION NEW AGENTS FOR TRANSFERRING NUCLEIC ACIDS, COMPOSITIONS CONTAINING THEM AND THEIR USES APPLICANT(S).FOR DO/EO/US Gerardo BYK, Marc FREDERIC, Hans HOFLAND, Daniel SCHERMAN Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information: This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. 2 This is a SECOND or SUBSEOUENT submission of items concerning a filing under 35 U.S.C. 371. This is an express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay 3. examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(1). A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date. A copy of the International Application as filed (35 U.S.C. 371(c)(2)) is transmitted herewith (required only if not transmitted by the International Bureau). has been transmitted by the International Bureau. is not required, as the application was filed in the United States Receiving Office (RO/US) A translation of the International Application into English (35 U.S.C. 371(c)(2)). Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)) are transmitted herewith (required only if not transmitted by the International Bureau). ь have been transmitted by the International Bureau. have not been made; however, the time limit for making such amendments has NOT expired. c. have not been made and will not be made. 8 A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)), and Power of Attorney. 10 A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)). Items 11, to 16, below concern other document(s) or information included: An information Disclosure Statement under 37 CFR 1.97 and 1.98. 12 An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. 13 A FIRST preliminary amendment A SECOND or SUBSEQUENT preliminary amendment. 14 A substitute specification. A change of power of attorney and/or address letter. Other items or information: 16 CERTIFICATION UNDER 37 CFR 1.10 "Express Mail" Mailing Number # 4685 154182 U.S Date of Deposit Oct. 2, 2000 I hereby certify that this paper (along with any referred to as being attached or enclosed) is being deposited with the United States Postal Service "Express Mail Post Office to Addressee" service under 37 CFR 1.10 on the date indicated

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Kelna L. Wetherill
(Signature of person mailing paper)

Debra L. Wetherill

(Type or print name of person mailing paper)

THE ADDITIONAL NO.	INTERNATIONAL A	APPLICATION NO	ATTORNEY'S DOC	KET NUMBER	
U.S. APPLICATION NO 7678					
(2.12.0.00.9.7	PCT/FR99/007	40	ST98009		
17. X The following fees are submitted: Basic National Fee (37 CFR 1.492(a)(1)-(5): Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445 (a)(2) paid to USPTO and International Search Report not prepared by the EPO or JPO\$2,000			CALCULATIONS	PTO use only	
International preliminary examination fee paid to USPTO (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO					
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International preliminary examination fee paid to USPTO (37 CFR 1.482) but all claims did not satisfy provisions of PCT Article 33(1)-(4)\$690					
International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(1)-(4)					
ENTER API	PROPRIATE BASIC	FEE AMOUNT =	\$860.00		
Surcharge of \$130.00 for furnishing the oath or declaration later than [] 20 [] 30 months from the earliest claimed priority date (37 CFR 1.492(c)).			\$		
Claims Number Filed	Number Extra	Rate	* 1000 00		
Total Claims 80 - 20 =	60	X \$ 18.00	\$ 1080.00 \$ 80.00		
Independent Claims 4 - 3 =	1	X \$ 80.00			
Multiple dependent claim(s) (if applicable)		+ 270.00	\$ 270.00 \$2290.00		
TOTAL OF ABOVE CALCULATIONS =					
Reduction by 1/2 for filing by small entity, if applicable. Verified Small Entity statement must also be filed. (Note 37 CFR 1.9, 1.27, 1.28).			S		
SUBTOTAL =			\$		
Processing fee of \$130.00 for furnishing the English translation later the [] 20 [] 30 months from the earliest claimed priority date (37 CFR 1.492(f)).			s		
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Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be					
accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property +			s		
TOTAL FEES ENCLOSED =			\$2290.00		
			Amount to be		
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a. A check in the amount of \$ to cover the above fee is enclosed. b. X Please charge my Deposit Account No. 18-1982 in the amount of \$_2290.00 to cover the above fees. A duplicate copy of this sheet is enclosed.					
c. X The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 18-1982. A duplicate copy of this sheet is enclosed.					
NOTE: Where an appropriate time limit und must be filed and granted to restore the appl	der 37 CFR 1.494 or 1.4 ication to pending status	195 has not been met, a p s.	etition to revive (37 Cl	FR 1.137(a) or (b))	
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ST98009

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NEW AGENTS FOR TRANSFERRING NUCLEIC ACIDS, COMPOSITIONS CONTAINING THEM AND THEIR USES

The present invention relates to new

5 compounds which are useful as agents for transferring nucleic acids into cells. These new compounds are more particularly related to the lipopolyamine family and comprise at least one cyclic amidine function. They are useful for transfecting nucleic acids to various types

10 of cells, in vitro, ex vivo or in vivo.

With the development of biotechnology, the possibility of effectively transferring nucleic acids into cells has become a necessity. It may involve the transfer of nucleic acids into cells in vitro, for 15 example for the production of recombinant proteins, or in the laboratory for studying the regulation of the expression of genes, the cloning of genes, or any other manipulation involving DNA. It may also involve the transfer of nucleic acids into cells in vivo, for 20 example for the creation of transgenic animals, the production of vaccines, carrying out labelling studies or also therapeutic approaches. It may also involve the transfer of nucleic acids into cells ex vivo, into approaches for bone marrow grafts, immunotherapy or 25 other methods involving the transfer of genes into cells collected from an organism for the purpose of their subsequent readministration.

Various types of synthetic vectors have been developed in order to improve the transfer of nucleic acid into cells. Among these vectors, cationic lipids possess advantageous properties. These vectors consist 5 of a cationic polar portion, which interacts with the nucleic acids, and of a hydrophobic lipid portion which favours cell penetration. Specific examples of cationic lipids are in particular the monocationic lipids (DOTMA : Lipofectin®), some cationic detergents (DDAB), lipopolyamines and in particular dioctadecylamidoglycyl spermine (DOGS) or palmitoylphosphatidylethanolamine 5-carboxyspermylamide (DPPES), whose preparation has been described, for example, in patent application EP 394,111. Another lipopolyamine family is represented 15 by the compounds as described in patent application WO 97/18185 incorporated into the present application by way of reference, and are illustrated in Figure 1.

However, up until now, injections into tissues, in particular the muscles, were often made 20 with non-formulated DNA in order to facilitate its entry into the cells, the combination with synthetic vectors leading to complexes which are too large in size to be incorporated into the cells.

It is one of the main problems which the

25 present invention proposes to solve. Indeed, compounds
according to the invention possess the unexpected
advantage of having a level of transfection in vivo
into the muscle which is at least equivalent to that

obtained with non-formulated DNA and in any case a very good level of transfection into the other tissues. The combination with a compound according to the invention protects the DNA from degradations by nucleases and/or 5 from deteriorations during freeze-drying, which contributes towards significantly improving the stability of the nucleolipid formulations. Furthermore, such a combination allows a slow controlled release of the nucleic acids.

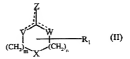
Moreover, the compounds according to the present invention belong to the family of cationic lipids and carry a novel cationic region which confers improved properties on said compounds, in particular reduced cytotoxicity compared with the prior art 15 cationic vectors. This cationic portion is indeed more precisely represented by one or more particular polyamine(s), carrying one or more cyclic amidine functions which very probably have the effect of "delocalizing" the positive charges, making the 20 compound less cationic overall, with the resulting beneficial effects known the toxicity point of view.

Thus, a first subject of the invention relates to new compounds, in D, L or DL form, of general formula (I):

for which:

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① CA represents a cycloamidine group and its mesomeric forms of general formula (II):



- 5 for which:
 - m and n are integers, independent of each other, of between 0 and 3 inclusive and such that m+n is greater than or equal to 1,
 - \bullet R_{1} represents a group of general formula (III):

$$-\frac{1}{2}(CH_2)_{p}-Y$$
 (III)

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for which p and q are integers, independent of each other, of between 0 and 10 inclusive, Y represents a carbonyl, amino, methylamino or methylene group, it being possible for Y to have different meanings within the different groups [(CH₂)_p-Y], and (*) represents either a hydrogen atom or is the site for bonding to the group Rep,

it being understood that R_1 may be bonded to any atom of general formula (II), including Z, and that there is a 20 single group R_1 in formula (II),

• X represents a group NR_2 or CHR_2 , R_2 being either a hydrogen atom or the bonding to the group R_1 as defined above,

- The group V represents:
- *1st case: a group of general formula (IV):

$$\begin{array}{ccc}
& & \text{NH} \\
& & & \\
R''N & & & \\
& & & & \\
\end{array}$$
(IV)

for which W' represents CHR \square or NR \square , and R" and R \square represent, independently of each other, a hydrogen atom, a methyl, or the bonding to the group R $_1$ as defined above, or

*2nd case: a group of general formula (V):

- for which W' represents CHRO or NRO, and R' and RO represent, independently of each other, a hydrogen atom, a methyl or the bonding to the group R1 as defined above.
 - @ Rep is absent or is a spacer of general
- 15 formula (VI):

$$- \left(\begin{array}{c} O \\ O \\ O \\ O \\ O \end{array} \right)$$

$$\left(\begin{array}{c} O \\ O \\ O \end{array} \right)$$

$$\left(\begin{array}{c} O \\ O \\ O \end{array} \right)$$

$$\left(\begin{array}{c} O \\ O \\ O \end{array} \right)$$

whose nitrogen atom is attached to the atoms X, V, W or Z or to the substituent Y of the group R_1 depending on the cases, and

 \circ t is an integer between 0 and 8 inclusive,

- r is an integer between 0 and 10 inclusive, it being possible for r to have different meanings within the different groups -NR₄-(CH) $_r$ -,
- R₃, which may have different meanings within the
 5 different groups NR₄-(CH)_rR₃, represents a hydrogen atom, a methyl group or a group of general formula (VII):

$$\frac{-\left(CH_{2}\right)_{5}-N}{R_{4}}H$$
 (VII)

for which u is an integer between 1 and 10 inclusive, s

10 is an integer between 2 and 8 inclusive which may have
different meanings within the different groups
-(CH₂)_s-NR₅, and R₅ is a hydrogen atom, a group CA as
defined above, it being understood that the groups CA
are independent from each other and may be different,

15 or a group of general formula (VII), it being
understood that the groups of general formula (VII) are
independent of each other and may have different
meanings,

- ullet R₄ is defined in the same manner as R₃ or represents a 20 group CA as defined above, it being understood that the groups CA are independent of each other and may be different, and
- ③ R is bonded to the carbonyl function of the group Rep of general formula (VI), or if Rep is absent, 25 R is bonded directly to the group CA, and represents:

1.0

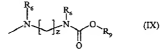
* either a group of formula NR_6R_7 for which R_6 and R_7 represent, independently of each other, a hydrogen atom or an optionally fluorinated, linear or branched, saturated or unsaturated aliphatic radical containing 1 to 22 carbon atoms, with at least one of the two substituents R_6 or R_7 different from hydrogen and the other containing between 10 and 22 carbon atoms,

* or a steroid derivative,

* or a group of general formula (VIII):

$$- \left[NH - - (CH_2)_x \right]_y - Q \qquad \quad (VIII)$$

for which x is an integer between 1 and 8 inclusive, y is an integer between 1 and 10 inclusive, and either Q represents a group $C(O)NR_6R_7$ for which R_6 and R_7 are as defined above, or Q represents a group $C(O)R_8$ for which R_8 represents a group of formula (IX):



for which z is an integer between 2 and 8 inclusive, and R₉ is an optionally fluorinated, saturated or unsaturated aliphatic radical containing 8 to 22 carbon 20 atoms, or a steroid derivative, and the two substituents R₆ are, independently of each other, as defined above, or R₈ represents a group -O-R₉ for which R₉ is as defined above.

According to one variant of the invention, the group R_1 is bonded either to Z or to V, on the one hand, and to the group Rep, on the other hand, via Y.

Advantageously, the cycloamidine group CA of 5 formula (II) comprises 5, 6, 7 or 8 members.

Moreover, in another variant of the invention, Rep is a spacer with 1, 2 or 3 "arms". The following spacers may for example be mentioned:

According to a second variant of the invention, R_3 represents a hydrogen atom or a methyl and R_4 is as defined above, or R_3 and R_4 present in

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formula (VI) represent hydrogen atoms, or R_4 is a hydrogen atom and R_3 is a group of formula (VII) in which R_5 represents a group CA.

Preferably, in formula (V), p and q are 5 chosen, independently of each other, from 2, 3 or 4.

In general, the group R contains at least one hydrophobic segment. For the purposes of the invention, "hydrophobic segment" is understood to mean any group of the lipid type, which promotes cell penetration. In particular, the group R contains at least one aliphatic chain or at least one steroid derivative.

According to a preferred variant, the group R represents a group of formula NR_6R_7 , R_6 and R_7 being as defined above, or represents a group of general 15 formula (VIII) in which Q represents a group $C(0)NR_6R_7$,

 R_6 and R_7 being as defined above.

Preferably, R_6 and/or R_7 represent, independently of each other, a saturated or unsaturated linear aliphatic chain containing 10 to 22 carbon

20 atoms, preferably with 12, 14, 16, 17, 18 or 19 carbon atoms. They are, for example, $(CH_2)_{11}CH_3$, $(CH_2)_{13}CH_3$, $(CH_2)_{15}CH_3$, $(CH_2)_{17}CH_3$ or oleyl groups and the like.

In a specific embodiment, the groups R_6 and R_7 are identical or different and each represent an 25 optionally fluorinated, linear or branched, saturated or unsaturated aliphatic chain containing 10 to 22 carbon atoms, as defined in the preceding paragraph.

When R represents a steroid derivative, the latter is advantageously chosen from cholesterol, cholestanol, $3-\alpha-5$ -cyclo- $5-\alpha$ -cholestan- $6-\beta$ -ol, cholic acid, cholesteryl formate, chotestanyl formate,

5 3α,5-cyclo-5α-cholestan-6β-yl formate, cholesterylamine, 6-(1,5-dimethylhexyl)-3a,5a-dimethylhexadecahydrocyclopenta[a]cyclopropa[2,3]cyclopenta[1,2-f]naphthalen-10-ylamine or cholestanylamine.

These new compounds of general formula (I)

- may be provided in the form of nontoxic and pharmaceutically acceptable salts. These nontoxic salts comprise salts with inorganic acids (hydrochloric, sulphuric, hydrobromic, phosphoric or nitric acids) or with organic acids (acetic, propionic, succinic,
- maleic, hydroxymaleic, benzoic, fumaric,
 methanesulphonic or oxalic acids) or with inorganic
 bases (sodium hydroxide, potassium hydroxide, lithium
 hydroxide or calcium hydroxide) or with organic bases
 (tertiary amines such as triethylamine, piperidine or
 benzylamine).

By way of example illustrating the preferred compounds according to the invention, the compounds of the following formulae may be mentioned:

Compound (1)

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N-dioctadecylcarbamoylmethyl-2-{3-[4-(2-iminotetrahydropyrimidin-1-yl)butylamino]propylamino}acetamide

Compound (2)

5 N-ditetradecylcarbamoylmethyl-2-{3-[4-(2-iminotetra-hydropyrimidin-1-yl)butylamino]propylamino}acetamide

Compound (3)

 $\label{eq:continuous} 2-(3-\{4-[3-(4,5-dihydro-1H-imidazol-2-ylamino)-propylamino] butylamino}-N-ditetradecylcarbamoylmethylacetamide$

Compound (4)

2-(3-{bis[3-(4,5-dihydro-1H-imidazol-2-ylamino)propyl]amino}propylamino)-N-ditetradecylcarbamoylmethylacetamide

Compound (5)

N-ditetradecylcarbamoylmethyl-2-{3-[3-(1,4,5,6-tetrahydropyrimidin-2-ylamino)propylamino)propylamino}acetamide

Compound (6)

N-dioctadecylcarbamoylmethyl-2-{3-[3-(1,4,5,6-tetrahydropyrimidin-2-ylamino)propylamino}acetamide

The compounds of the invention may be 10 prepared in various ways. According to a first method, the compounds of the invention may be obtained by synthesis of analogous lipopolyamines (that is to say the same structure but with no cycloamidine group), the cyclization to cycloamidine groups being carried out in 15 a second instance. The analogous lipopolyamines may be obtained by any method known to persons skilled in the art, and in particular according to the methods described in Application WO 97/18185 or by similar methods. The cyclization of the amidine heads may, for 20 example, be carried out by reaction between one and/or more primary amines of the lipopolyamine and reagents such as O-methylisourea sulphate hydrogen sulphate [J. Med. Chem., 1995, 38(16), pp. 3053-3061] or S-methylisothiourea hemisulphate [Int. J. Pept. Prot. 25 Res., 1992, 40, pp. 119-126]. Preferably, the procedure

is carried out in aqueous medium in the presence of a

base in the hot state [J. Med. Chem. 1985, pp. 694-698 and J. Med. Chem., 1996, pp. 669-672]. As preferred solvents, there may be mentioned water/alcohol mixtures or dimethylformamide. As base, triethylamine,

5 N-ethyldiisopropylamine, sodium hydroxide, potassium hydroxide and the like may be used. The temperature is preferably between 40°C and 60°C, and still more preferably the reaction is carried out at 50°C.

Another method consists in carrying out a

10 synthesis of building blocks carrying the cycloamidine
function which are then grafted onto lipids equipped
with spacers. This method has the advantage of
providing access to a large number of products. For the
purposes of the invention, "blocks" is understood to

15 mean any functional segment of the molecule. For
example, the cycloamidine group CA as defined in
general formula (II), Rep or R each constitute distinct

By way of example, the procedure may, for 20 example, be carried out in the following manner:

blocks for the purposes of the invention.

1) Synthesis of the building block R:

- a) When R represents -NR₆R₇, either it is commercially available, or it can be synthesized according to one of the following methods:
- by alkylative reduction between an amine carrying the group R_6 and an aldehyde carrying the group R_7 . The procedure is preferably carried out in a chlorinated solvent (for example dichloromethane, chloroform,

1,2-dichloroethane and the like [J. Org. Chem., 1996, pp. 3849-3862]) or in any other organic solvent which is compatible with the reaction (for example tetrahydrofuran), in the presence of sodium

5 triacetoxyborohydride, sodium cyanoborohydride or derivatives thereof (for example lithium cyanoborohydride) [J. Am. Chem. Soc., 1971, pp. 2897-

29041 and acetic acid.

- or by substitution of a leaving group carried by R₆,
 by an amine carrying the group R₇. By way of example of a leaving group, halogen atoms (Br, Cl, I) or tosyl or mesyl substituents and the like may be mentioned. The procedure is preferably carried out in the presence of a basic reagent, for example sodium carbonate,
- 15 potassium hydroxide, sodium hydroxide, triethylamine and the like, in an alcohol (for example ethanol) under reflux [J. Am. Chem. Soc., 1996, pp. 8524-8530].
 - or by coupling between a fatty acid (or derivatives thereof such as fatty acid chlorides) and a fatty
- amine. The amide obtained is then reduced by a hydride, for example lithium aluminium hydride or any other hydride known to persons skilled in the art, in an ether (for example tetrahydrofuran (THF), t-butyl methyl ether (TBME), dimethoxyethane (DME) and the
- 25 like).
 - b) When R represents a group of general formula (VIII), the peptide coupling is carried out between the group Q and H-[NH-(CH₂) $_{\rm x}$] $_{\rm y}$ COOH. The peptide coupling is

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carried out according to conventional methods known to persons skilled in the art (Bodanski M., Principles and Practices of peptide Synthesis, Ed. Springe-Verlag) or by any known similar method. In particular, the reaction may be carried out in the presence of a nonnucleophilic base in suitable aprotic solvents (such as chloroform, dimethylformamide, methylpyrrolidone, acetonitrile, dichloromethane and the like), at a temperature of between 0 and 100°C, the pH being adjusted between 9 and 11.

Q is either commercially available or, when O represents a group C(O)R8 with R8 of formula (IX), it may be synthesized by reaction between a chloroformate 15 which is commercially available (for example cholesteryl chloroformate) or obtained according to conventional methods known to persons skilled in the art from a commercially available chloroformate, and a diamine which is commercially available (for example 20 N-ethylenediamine) or obtained according to conventional methods known to persons skilled in the art. Preferably, the procedure is carried out in a chlorinated solvent (for example dichloromethane, chloroform, 1,2-dichloroethane and the like) or in any 25 other organic solvent which is compatible with the reaction, such as for example dimethylformamide, dimethyl sulphoxide, acetonitrile and the like.

The group H-[NH-(CH2)x]y-COOH is a commercially available amino acid when y is equal to 1, or is obtained by one or more cyanoethylation reactions according to the method described below in the 5 synthesis of Rep when y is greater than 1.

2) Synthesis of the building block Rep:

The group Rep is obtained by cyanoethylation or by dicyanoethylation (depending on whether it is desired to obtain a linear or branched Rep structure) of an amino acid of formula HOOC-(CH₂)_r-NH₂ and then by reduction of the nitrile functions into amines.

a) Mono- or dicyanoethylation:

$$\frac{\text{HOOC-(CH2)r-NH2} + 1 \text{ or } 2}{R'}$$
or
$$R'-N$$

$$R'-N$$

$$R'-N$$

$$R'-N$$

$$r-1$$

$$CN$$

Preferably, the procedure is carried out in a basic

15 aqueous medium. For example, the reaction is carried out in solvents such as water, alcohols (for example methanol, ethanol and the like), in the presence of a base such as sodium hydroxide, potassium hydroxide, triethylamine and the like. In the case of

20 monocyanoethylation, the work is preferably carried out

in the cold state [J. Am. Chem. Soc., 1950, pp. 2599-2603]. In the case of dicyanoethylation, the work is preferably carried out in the hot state and with an excess of acrylonitrile [J. Am. Chem. Soc., 1951, 5 pp. 1641-1644].

b) The reduction of the nitrile functions into amines is carried out by catalytic hydrogenation in a basic medium or by any other method known to a person skilled in the art. By way of example, it is possible to use
platinum oxide or Raney nickel [J. Org. Chem., 1988, pp. 3108-3111] as catalyst. Preferably, the solvent chosen is an alcohol (for example methanol, ethanol and

the like) in the presence of a base, for example sodium

hydroxide, potassium hydroxide and the like.

3) Synthesis of the building block Rep-R:

The building block Rep-R is obtained by peptide coupling between the acid Rep and the amine R which are obtained in the preceding steps.

The peptide coupling is carried out according to

20 conventional methods known to persons skilled in the
art (Bodanski M., Principles and Practices of peptide

Synthesis, Ed. Springe-Verlag) or by any known similar
method. In particular, the reaction may be carried out
in the presence of a nonnucleophilic base, in suitable

25 aprotic solvents (such as chloroform.

dimethylformamide, methylpyrrolidione, acetonitrile, dichloromethane and the like), at a temperature of

between 0 and 100°C, the pH being adjusted between 9 and 11.

- Synthesis of the compounds according to the invention CA-Rep-R:
- 5 The compounds according to the invention are obtained according to several possible methods:
 - a) By coupling in a basic medium between the terminal amine present on Rep-R obtained in the preceding step, and CA-S-CH₃, according to conventional methods known to
- 10 persons skilled in the art. The procedure is preferably carried out in a chlorinated solvent (for example dichloromethane, chloroform and the like) or in other organic solvents compatible with the reaction, such as for example water, alcohols, dimethylformamide and the
- 15 like, in the presence of a base (for example triethylamine, sodium hydroxide, potassium hydroxide, N-ethyldiisopropylamine and the like), and at room temperature (about 20°C).

The building block ${\rm CA-S-CH_3}$ is either commercially

- 20 available (that is the case for example for 2-methylthio-2-imidazoline hydriodide), or it can be obtained by the action of a carbon disulphide on an appropriate diamine (that is to say chosen as a function of the cycloamidine group which it is desired
- 25 to obtain), followed by a methylation. For example, the reaction scheme may be illustrated in the following manner:

Preferably, the reaction process is carried out in an

alcohol (for example ethanol). The methylation step is carried out by the action of a halomethyl, it being 5 possible for the halogen atom to be, for example, an iodine atom [J. Am. Cem. Soc., 1956, pp. 1618-1620 and Bioorg. Med. Chem. Lett., 1994, pp. 351-354]. b) By internal cyclization of the cycloamidine group from the amino functions present on Rep-R, by the 10 action of O-methylisourea hydrogen sulphate or Smethylisothiourea hemisulphate. Preferably, the procedure is carried out in an aqueous medium in the presence of a base in the hot state [J. Med. Chem., 1985, pp. 694-698 and J. Med. Chem., 1996, pp. 669-15 672]. As a preferred solvent, water/alcohol mixtures or dimethylformamide may be mentioned. As a base, triethylamine, N-ethyldiisopropylamine, sodium hydroxide, potassium hydroxide and the like may be used. The temperature is preferably between 40°C and 20 60°C, and still more preferably the reaction is carried out at 50°C.

c) By peptide coupling between CA-COOH and Rep-R according to conventional techniques known to persons skilled in the art, as described above.

The building block CA-COOH may be obtained in various 5 ways:

- ullet by the action of a building block CA-S-CH $_3$ on an amino acid or a polyamino acid according to methods known to persons skilled in the art or by any other similar method [J. Am. Chem. Soc., 1956,
- 10 pp. 1618-1620]. The building block CA-S-CH₃ is obtained in the same manner as above, and the amino or polyamino acid is chosen as a function of the desired compound according to the invention, or
- by the action of an S,S-dimethyltosylimino thiocarbonimidate or of one of its derivatives on a polyamino acid according to methods known to persons skilled in the art or by any similar method [J. Org. Chem., 1971, pp. 46-48]. Preferably, the procedure is carried out in an ethanolic medium in the presence of a
 base (for example sodium hydroxide) and at the reflux
 - By way of example of building blocks CA-COOH which may be obtained by one of the methods described above, the

following building blocks may be mentioned:

temperature of the mixture.

In all the reactions disclosed above, when
the amino substituents present in the various groups
may interfere with the reactions carried out, it is

5 preferable to protect them beforehand with compatible
radicals which can be introduced and removed without
affecting the rest of the molecule. By way of example,
the protective radicals may be chosen from the radicals
described by T.W. GREENE, Protective Groups in Organic

10 Synthesis, J. Wiley-Interscience Publication (1991) or
by McOmie, Protective Groups in Organic Chemistry,
Plenum Press (1973).

Another subject of the invention relates to a composition comprising at least one compound of formula

15 (I) as defined above. In particular, another subject according to the present invention comprises a compound of formula (I) as defined above and a nucleic acid.

When a compound according to the invention and a nucleic acid are brought into contact, they form a complexes by interaction between the positive charges present at physiological pH on the compound according 5 to the invention and the negative charges of the nucleic acid. This complex is called "nucleolipid complex" in the remainder of the text which follows. Preferably, the compound according to the invention and the nucleic acid are present in quantities such that 10 the ratio of the positive charges of the compound to the negative charges of the nucleic acid is between 0.1 and 50, preferably between 0.1 and 20. This ratio can be easily adjusted by persons skilled in the art according to the compound used, the nucleic acid and 15 the desired applications (in particular the type of cells to be transfected).

For the purposes of the invention, "nucleic acid" is understood to mean both a deoxyribonucleic acid and a ribonucleic acid. They may be natural or

20 artificial sequences, and in particular genomic DNA
(gDNA), complementary DNA (cDNA), messenger RNA (mRNA),
transfer RNA (tRNA), ribosomal RNA (rRNA), hybrid
sequences or synthetic or semisynthetic sequences,
oligonucleotides which are modified or otherwise. These
25 nucleic acids may be of human, animal, plant, bacterial
or viral origin and the like. They may be obtained by
any technique known to persons skilled in the art, and
in particular by the screening of libraries, by

chemical synthesis or by mixed methods including the chemical or enzymatic modification of sequences obtained by the screening of libraries. They may be chemically modified.

As regards more particularly deoxyribonucleic acids, they may be single- or double-stranded, as well as short oligonucleotides or longer sequences. In particular, the nucleic acids advantageously consist of plasmids, vectors, episomes, expression cassettes and 10 the like. These deoxyribonucleic acids may carry a replication origin which is functional or otherwise in the target cell, one or more marker genes, sequences for regulating transcription or replication, genes of therapeutic interest, anti-sense sequences which are 15 modified or otherwise, regions for binding to other cellular components, and the like.

Preferably, the nucleic acid comprises an expression cassette consisting of one or more genes of therapeutic interest under the control of one or more 20 promoters and a transcriptional terminator which are active in the target cells.

For the purposes of the invention, "cassette for expression of a gene of interest" is understood to mean a DNA fragment which may be inserted into a vector 25 at specific restriction sites. The DNA fragment comprises a nucleic acid sequence encoding an RNA or a polypeptide of interest and comprises, in addition, the sequences necessary for the expression (enhancer(s),

promoter(s), polyadenylation sequences and the like) of said sequence. The cassette and the restriction sites are designed to ensure insertion of the expression cassette into a reading frame appropriate for transcription and translation.

It is generally a plasmid or an episome carrying one or more genes of therapeutic interest. By way of example, there may be mentioned the plasmids described in patent applications WO 96/26270 and

10 WO 97/10343 which are incorporated into the present application by way of reference.

For the purposes of the invention, gene of therapeutic interest is understood to mean in particular any gene encoding a protein product having a 15 therapeutic effect. The protein product thus encoded may in particular be a protein or a peptide. This protein product may be exogenous, homologous or endogenous in relation to the target cell, that is to say a product which is normally expressed in the target 20 cell when the latter has no pathological condition. In this case, the expression of a protein makes it possible, for example, to palliate an insufficient expression in the cell or the expression of a protein which is inactive or weakly active because of a 25 modification, or to overexpress said protein. The gene of therapeutic interest may also encode a mutant of a cellular protein, having increased stability, a modified activity and the like. The protein product may also be heterologous in relation to the target cell. In this case, an expressed protein may, for example, supplement or provide an activity which is deficient in the cell, allowing it to combat a pathological condition, or to stimulate an immune response.

Among the therapeutic products for the

purposes of the present invention, there may be mentioned more particularly enzymes, blood derivatives, hormones, lymphokines: interleukins, interferons, TNF, and the like (FR 92/03120), growth factors, neurotransmitters or their precursors or synthesis enzymes, trophic factors (BDNF, CNTF, NGF, IGF, GMF, aFGF, bFGF, NT3, NT5, HARP/pleiotrophin and the like), apolipoproteins (ApoAI, ApoAIV, ApoE, and the like, FR 93/05125), dystrophin or a minidystrophin (FR 91/11947), the CFTR protein associated with cystic fibrosis, tumour suppressor genes (p53, Rb, Rap1A, DCC,

20 IX), the genes involved in DNA repair, suicide genes (thymidine kinase, cytosine deaminase), the genes for haemoglobin or other protein carriers, metabolic enzymes, catabolic enzymes and the like.

k-rev, and the like, FR 93/04745), the genes encoding factors involved in coaquilation (factors VII, VIII,

The nucleic acid of therapeutic interest may

25 also be a gene or an anti-sense sequence, whose
expression in the target cell makes it possible to
control the expression of genes or the transcription of
cellular mRNAs. Such sequences can, for example, be

transcribed in the target cell into RNAs which are complementary to cellular mRNAs and thus block their translation to protein, according to the technique described in Patent EP 140 308. The therapeutic genes also comprise the sequences encoding ribozymes, which are capable of selectively destroying target RNAs (EP 321 201).

As indicated above, the nucleic acid may also comprise one or more genes encoding an antigenic

10 peptide, which is capable of generating an immune response in humans or in animals. In this specific embodiment, the invention allows either the production of vaccines or the carrying out of immunotherapeutic treatments applied to humans or to animals, in

15 particular against microorganisms, viruses or cancers. They may be in particular antigenic peptides specific for the Epstein-Barr virus, the HIV virus, the hepatitis B virus (EP 185 573), the pseudo-rabies virus, the syncitia forming virus, other viruses, or

20 antigenic peptides specific for tumours (EP 259 212).

Preferably, the nucleic acid also comprises sequences allowing the expression of the gene of therapeutic interest and/or the gene encoding the antigenic peptide in the desired cell or organ. They

25 may be sequences which are naturally responsible for the expression of the gene considered when these sequences are capable of functioning in the infected cell. They may also be sequences of different origin

(responsible for the expression of other proteins, or even synthetic). In particular, they may be promoter sequences of eukaryotic or viral genes. For example, they may be promoter sequences derived from the genome of the cell which it is desired to infect. Likewise, they may be promoter sequences derived from the genome of a virus. In this regard, there may be mentioned, for example, the promoters of the E1A, MLP, CMV and RSV genes, and the like. In addition, these expression
sequences may be modified by the addition of activating or regulatory sequences, and the like. The promoter may also be inducible or repressible.

Moreover, the nucleic acid may also comprise, in particular upstream of the gene of therapeutic

15 interest, a signal sequence directing the therapeutic product synthesized in the secretory pathways of the target cell. This signal sequence may be the natural signal sequence of the therapeutic product, but it may also be any other functional signal sequence, or an artificial signal sequence. The nucleic acid may also comprise a signal sequence directing the synthesized therapeutic product toward a particular compartment of the cell.

The compositions according to the invention

25 may, in addition, comprise one or more adjuvants

capable of combining with the complexes formed between

the compound according to the invention and the nucleic

acid, and of improving the transfecting power thereof.

In another embodiment, the present invention therefore relates to compositions comprising a nucleic acid, a compound of formula (I) as defined above and one or more adjuvants capable of combining with the compound 5 (I)/nucleic acid nucleolipid complexes and of improving the transfecting power thereof. The presence of this type of adjuvants (for example lipids, peptides or proteins) can advantageously make it possible to increase the transfecting power of the compounds.

In this regard, the compositions of the invention may comprise, as adjuvant, one or more neutral lipids. Such compositions are particularly advantageous, in particular when the charge ratio R is low. The applicant has indeed shown that the addition 15 of a neutral lipid makes it possible to improve the formation of the nucleolipid particles and to promote the penetration of the particle into the cell by destabilizing its membrane.

More preferably, the neutral lipids used 20 within the framework of the present invention are lipids containing two fatty chains. In a particularly advantageous manner, natural or synthetic lipids which are zwitterionic or lacking ionic charge under physiological conditions are used. They may be chosen 25 more particularly from dioleoylphosphatidylethanolamine (DOPE), oleoylpalmitoylphosphatidylethanolamine (POPE), di-stearoyl, -palmitoyl, -mirystoylphosphatidylethanolamines as well as their derivatives which are

N-methylated 1 to 3 times, phosphatidylglycerols, diacylglycerols, glycosyldiacylglycerols, cerebrosides (such as in particular galactocerebrosides), sphingolipids (such as in particular sphingomyelins) or asialogangliosides (such as in particular asialoGM1 and GM2).

These different lipids may be obtained either by synthesis or by extraction from organs (for example the brain) or from eggs, by conventional techniques

10 well known to persons skilled in the art. In particular, the extraction of the natural lipids may be carried out by means of organic solvents (see also Lehninger, Biochemistry).

More recently, the applicant has demonstrated

that it was also particularly advantageous to use, as
adjuvant, a compound involved directly or otherwise in
the condensation of the nucleic acid (WO 96/25508). The
presence of such a product in a composition according
to the invention makes it possible to reduce the

quantity of compound of formula (I), with the
beneficial consequences resulting therefrom from the
toxicological point of view, without any damaging
effect on the transfecting activity. Product involved
in the condensation of the nucleic acid is intended to
define a product which compacts, directly or otherwise,
the nucleic acid. More precisely, this product may
either act directly at the level of the nucleic acid to
be transfected, or may be involved at the level of an

additional product which is directly involved in the condensation of this nucleic acid. Preferably, it acts directly at the level of the nucleic acid. For example, the precompacting agent may be any polycation, for 5 example polylysine. According to a preferred embodiment, this product which is involved in the condensation of the nucleic acid is derived as a whole or in part from a protamine, a histone or a nucleolin and/or from one of their derivatives. Such an agent may 10 also consist, as a whole or in part, of peptide units (KTPKKAKKP) and/or (ATPAKKAA), it being possible for the number of units to vary between 2 and 10. In the structure of the compound according to the invention, these units may be repeated continuously or otherwise. 15 They may thus be separated by linkages of a biochemical nature, for example by one or more amino acids, or of a

Preferably, the compositions of the invention comprise from 0.01 to 20 equivalents of adjuvant(s) for 20 one equivalent of nucleic acids in mol/mol and, more preferably, from 0.5 to 5.

chemical nature.

In a particularly advantageous embodiment, the compositions according to the present invention comprise, in addition, a targeting element which makes it possible to orient the transfer of the nucleic acid. This targeting element may be an extracellular targeting element which makes it possible to orient the transfer of DNA toward certain cell types or certain

desired tissues (tumour cells, hepatic cells, haematopoietic cells and the like). It may also be an intracellular targeting element which makes it possible to orient the transfer of the nucleic acid toward 5 certain preferred cellular compartments (mitochondria, nucleus and the like). The targeting element may be linked to the compound according to the invention or also to the nucleic acid as specified above.

Among the targeting elements which may be 10 used within the framework of the invention, there may be mentioned sugars, peptides, proteins, oligonucleotides, lipids, neuromediators, hormones, vitamins or derivatives thereof. Preferably, they are sugars, peptides or proteins such as antibodies or 15 antibody fragments, ligands of cell receptors or fragments thereof, receptors or receptor fragments, and the like. In particular, they may be ligands of growth factor receptors, cytokine receptors, cellular lectintype receptors, or RGD sequence-containing ligands with 20 an affinity for the receptors for adhesion proteins such as the integrins. There may also be mentioned the receptors for transferrin, HDLs and LDLs, or the folate transporter. The targeting element may also be a sugar which makes it possible to target lectins such as the 25 receptors for asialoglycoproteins or for syalydes such as the sialyde Lewis X, or alternatively an Fab fragment of antibodies, or a single-chain antibody (ScFv).

The combination of the targeting elements with the nucleolipid complexes of the invention may be made by any technique known to persons skilled in the art, for example by coupling to a hydrophobic part or to a part which interacts with the nucleic acid of the compound of general formula (I) according to the invention, or alternatively to a group which interacts with the compound of general formula (I) according to the invention or with the nucleic acid. The

10 interactions in question may be, according to a preferred mode, of an ionic or covalent nature.

According to another variant, the compositions of the invention may also incorporate at least one nonionic surfactant in a sufficient quantity to stabilize the size of the particles of compound of general formula (I)/nucleic acid nucleolipid complexes. The introduction of nonionic surfactants prevents the formation of aggregates, which makes the composition more particularly suitable for an in vivo administration. The compositions according to the invention incorporating such surfactants have an advantage from the point of view of safety. They also have an additional advantage in the sense that they reduce the risk of interference with other proteins,

The surfactants advantageously consist of at least one hydrophobic segment, and at least one

compositions of nucleolipid complexes.

hydrophilic segment. Preferably, the hydrophobic segment is chosen from aliphatic chains, polyoxyalkylenes, alkylidene polyesters, polyethylene glycols with a benzyl polyether head and cholesterol, 5 and the hydrophilic segment is advantageously chosen from polyoxyalkylenes, polyvinyl alcohols, polyvinyl pyrrolidones or saccharides. Such nonionic surfactants have been described in application WO 98/34648.

The subject of the invention is also the use 10 of the compounds of general formula (I) as defined above to manufacture a medicament for treating diseases by transfer of nucleic acids (and more generally of polyanions) into primary cells or into established lines. They may be in particular fibroblast cells, 15 muscle cells, nerve cells (neurons, astrocytes, qlyal cells), hepatic cells, haematopoietic cell lines (lymphocytes, CD34, dendritic cells and the like), epithelial cells and the like, in differentiated or pluripotent form (precursors).

Such a use is particularly advantageous because the compounds of general formula (I) according to the invention have a reduced cytotoxicity compared with the prior art cationic lipids. The applicant has in particular demonstrated that at very high charge 25 ratios which normally result in the death of the animals following transfection, no apparent cytotoxicity was detected. The compounds of the invention may be used in particular for the in vitro,

ex vivo or in vivo transfection of nucleic acids. For uses in vivo, for example in therapy or for studying the regulation of genes or the creation of animal models of pathological conditions, the compositions 5 according to the invention can be formulated for administration by the topical, cutaneous, oral, rectal, vaginal, parenteral, intranasal, intravenous, intramuscular, subcutaneous, intraocular, transdermal, intratracheal or intraperitoneal route, and the like. 10 Preferably, the compositions of the invention contain a vehicle which is pharmaceutically acceptable for an injectable formulation, in particular for a direct injection into the desired organ, or for administration by the topical route (on the skin and/or the mucous 15 membrane). They may be in particular isotonic sterile solutions, or dry, in particular freeze-dried, compositions which, upon addition, depending on the case, of sterilized water or of physiological saline, allow the constitution of injectable solutions. The 20 nucleic acid doses used for the injection as well as the number of administrations may be adapted according to various parameters, and in particular according to the mode of administration used, the relevant pathological condition, the gene to be expressed, or 25 the desired duration of treatment. As regards more particularly the mode of administration, it may be either a direct injection into the tissues, for example

at the level of the tumours, or the circulatory system,

or a treatment of cells in culture followed by their reimplantation in vivo by injection or transplantation. The relevant tissues within the framework of the present invention are, for example, the muscles, skin, 5 brain, lungs, liver, spleen, bone marrow, thymus, heart, lymph, blood, bones, cartilages, pancreas, kidneys, bladder, stomach, intestines, testicles, ovaries, rectum, nervous system, eyes, glands, connective tissues, and the like. Advantageously, the transfected tissues are the muscles and the lungs.

The invention relates, in addition, to a method of transferring nucleic acids into cells comprising the following steps:

- (1) bringing the nucleic acid into contact with a 15 compound of general formula (I) as defined above, to form a nucleolipid complex, and
 - (2) bringing the cells into contact with the nucleolipid complex formed in (1).

The cells may be brought into contact with

20 the nucleolipid complex by incubating the cells with
said complex (for uses in vitro or ex vivo), or by
injecting the complex into an organism (for uses in
vivo). The incubation is preferably carried out in the
presence, for example, of 0.01 to 1000 µg of nucleic

25 acid per 10⁶ cells. For administration in vivo, nucleic
acid doses of between 0.01 and 10 mg may for example be
used.

In the case where the compositions of the invention contain, in addition, one or more adjuvants as defined above, the adjuvant(s) is (are) previously mixed with the compound of general formula (I) according to the invention or with the nucleic acid.

The present invention thus provides a particularly advantageous method for the treatment of diseases by administration of a nucleic acid encoding a protein or which can be transcribed into a nucleic acid capable of correcting said disease, said nucleic acid being combined with a compound of general formula (I) as defined above, under the conditions defined above. More particularly, this method is applicable to diseases resulting from a deficiency in a protein or nucleic product, the administered nucleic acid encoding said protein product or being transcribed into a nucleic product or constituting said nucleic product.

The invention extends to any use of a compound of formula (I) according to the invention for the in vivo, ex vivo or in vitro transfection of cells.

In addition to the preceding arrangements, the present invention also comprises other characteristics and advantages which will emerge from the examples and figures below, which should be considered as illustrating the invention without limiting its scope. In particular, the applicant proposes, with no limitation being implied, various operating protocols as well as reaction intermediates

which can be used to prepare the compounds of general formula (I). Of course, it is within the capability of persons skilled in the art to draw inspiration from these protocols and/or intermediate products in order to develop similar methods to lead to other compounds of general formula (I) according to the invention.

FIGURES

Figure 1: Structure of the synthetic vectors called

10 lipid A, lipid B, lipid c and lipid D in the present
invention and which are described in patent application
WO 97/18185 incorporated into the present application
by way of reference.

<u>Figure 2:</u> Schematic representation of the plasmid pxL2774.

Figure 3: Phase diagram for the compound (1)/DNA
nucleolipid complexes. The binding of compound (1) to
DNA was determined by following the decrease in the
fluorescence (in %, 100% being the fluorescence of the
20 naked DNA) of ethidium bromide (EtBr) (symbol ●, solid
line), as described according to the y-axis situated on
the right. The size of the particles of complexes
(in nm) is indicated on the y-axis situated on the
left. The x-axis represents the transfer agents/DNA
25 charge ratio. The size of the nucleolipid complexes
without co-lipid is represented by the symbol ■ as a
solid line. The size of the nucleolipid complexes
containing 25% cholesterol is represented by the symbol

□ as a discontinuous line. The size of the nucleolipid complexes containing 40% DOPE is represented by the symbol ◆ as a discontinuous line. The method does not make it possible to determine the size of the particles above 3 µm.

Figure 4: Activity for in vitro gene transfer into HeLa cells of the nucleolipid complexes containing compound

(1) according to the present invention without co-lipid (dark-shaded middle bar), with 25% cholesterol (medium-shaded left-hand bar), and with 40 mol % of DOPE (light-shaded right-hand bar), compared with naked DNA. Only the nucleolipid complexes in which the DNA is completely saturated with the compound according to the invention and whose size is between 100 nm and 300 nm

Figure 5: Gene transfer activity in vitro into HeLa cells of the nucleolipid complexes formed from compound (3). The expression of luciferase, expressed in pg per well transfected, is represented on the y-axis. The charge ratio between compound (3) and the DNA in nmol/µg is represented on the x-axis. The expression was measured each time for formulations without colipid (micelles), with DOPE and with cholesterol.

Figure 6: Gene transfer activity in vitro into HeLa cells of the nucleolipid complexes formed from compound (5). The expression of luciferase, expressed in pg per well transfected, is represented on the y-axis. The charge ratio between compound (5) and the DNA in

nmol/µg is represented on the x-axis. The expression was measured each time for formulations without colipid (micelles), with DOPE and with cholesterol. Figure 7: Gene transfer activity in vitro into HeLa 5 cells of the nucleolipid complexes formed from compound (6). The expression of luciferase, expressed in pg per well transfected, is represented on the y-axis. The charge ratio between compound (6) and the DNA in nmol/µg is represented on the x-axis. The expression 10 was measured each time for formulations without colipid (micelles), with DOPE and with cholesterol. Figure 8: Gene transfer activity in vivo after direct injection into the muscle of the complexes containing compound (1) according to the present invention or the 15 compound of formula $H_2N(CH_2)_3NH(CH_2)_4NH(CH_2)_3NHCH_2COArgN[(CH_2)_{17}CH_3]_2$ (called "lipid A" in the remainder of the text which follows) without co-lipid (dark-shaded bar), with 25% cholesterol (medium-shaded bar), and with 40 mol % of 20 DOPE (light-shaded bar), compared with naked DNA. Only the complexes in which the DNA is completely saturated with lipid and whose size is between 100 nm and 300 nm

Figure 9: The importance of the invention is

25 illustrated by comparing the gene transfer activity of
two different lipids, compound (1) according to the
invention and lipid A, and of naked DNA via two routes
of administration: by the intravenous (iv) route and by

were used.

the intramuscular (im) route. Only the complexes in which the DNA is completely saturated with lipid and whose size is between 100 nm and 300 nm were used.

Figure 10: Gene transfer activity in vivo 48 hours

after i.m. injection of the nucleolipid complexes containing compounds (5) or (6) according to the present invention without co-lipid and at a charge ratio of 0.25/1, compared with naked DNA. The expression is expressed in pg of luciferase per ml.

Starting from the left, the bars represent: (a)

MATERIALS AND METHODS

(d) compound (6).

15 A\ MATERIALS

art.

- The starting amino or polyamino acids (or derivatives thereof) are commercially available. This is the case, for example, for N-(3-aminopropyl)glycine, N-(2-cyanoethyl)glycine or 2,4-diaminobutyric acid, or may be synthesized by conventional methods known to persons skilled in the art.

negative control; (b) naked DNA; (c) compound (5) and

- The cyclic isothioureas are also commercially available products, such as for example 2-methylthio-2imidazoline hydriodide, or may be synthesized by 25 conventional methods known to persons skilled in the
 - The amines substituted with one or more lipid(s) are commercially available or are synthesized from the

corresponding amines and aldehydes by alkylative reduction.

- The products such as triethylamine, trifluoroacetic acid, benzotriazol-1-yloxytris(dimethylamino)-
- 5 phosphonium hexafluorophosphate (BOP),
 dimethylaminopyridine (DMAP), benzyl chloroformate,
 di-tert-butyl dicarbonate are commercially available
 products. The sodium chloride and sodium carnbonate
 solutions are saturated. The potassium sulphate
- 10 solution has a concentration of 0.5 M.

B\ METHODS

1) Physical measurements

The Proton NMR spectra were recorded on Bruker 400 and 600 MHZ spectrometers.

15 The mass spectra were taken on an API-MS/III.

Methods of purification and analysis

- a) Direct-phase chromatography conditions
- The thin-layer chromatographies (TLC) were carried out on 0.2 mm thick Merck silica gel plates.
- 20 They are developed either under U.V. (254 nm), with ninhydrin, by spraying (light spray) an ethanolic solution of ninhydrin (40 mg/100 cm³ of ethanol) in order to reveal the amines or the amides by heating to 150°C, with fluorescamine, by spraying a solution
- 25 (40 mg/100 cm³ of acetone) in order to reveal the primary amines, with bromocresol green, by spraying a solution (0.1% in 2-propanol) in order to reveal the acids, with vanillin by spraying (light spray) an

ethanolic solution of vanillin (3%) with 3% sulphuric acid followed by heating to 120°C, or with iodine by covering the plate with iodine powder.

- The column chromatographies were carried out on a
- 5 Merck 60 silica gel having a particle size of 0.063-0.200 mm.
 - b) Preparative HPLC (High-Performance Liquid Chromatography) purification conditions

The equipment is a set for liquid-phase chromatography

10 in gradient mode, allowing U.V. detection. This preparative chain is composed of the following components:

Pump A: GILSON model 305 equipped with a 50 SC head.Pump B: GILSON model 303 equipped with a 50 SC head.

15 Injection loop: 5 ml.

Pressure module: GILSON model 806.

<u>Mixer</u>: GILSON model 811 C equipped with a 23 ml head.
<u>UV detector</u>: GILSON model 119 equipped with a preparative cell.

20 <u>Fraction collector</u>: GILSON model 202 equipped with No. 21 racks and a 10 ml glass tube.

Integrator: SHIMADZU model C-R6A.

<u>Column</u>: Column C4 (10 mm) made of stainless steel 25 cm long and 2.2 cm in diameter, marketed by VYDAC model

25 214 TP 1022.

The solution of product to be purified is loaded onto the column by means of the injection loop, the eluent is recovered in fractions of one tube in 30 seconds. The detector is set at the wavelength of 220 nm and $254\ nm$.

the mobile phases are defined as follows:

Solvent A

Solvent B

Demineralized water $2500~\text{cm}^3$ Acetonitrile for HPLC $2500~\text{cm}^3$ Trifluoroacetic acid $2~\text{cm}^3$ Trifluoroacetic acid $2.5~\text{cm}^3$

Gradient:

Time in minutes	% of	% of	Flow rate
	solvent A	solvent B	in cm³/min
0	90	10	18
10	90	10	18
110	0	100	18
120	0	100	18

5 c) Analytical chromatography techniques

- The HPLC (High-Performance Liquid Chromatography) analyses were carried out on a Merck-Hitachi apparatus equipped with a HITACHI D 2500 integrator-calculator, an autosampler AS-2000A, an intelligent pump L-6200A,
- - The columns for the analytical separations are Browlee columns made of stainless steel 3 cm long and 0.46 cm in diameter, marketed by APPLIED BIOSYSTEM.
- 15 The stationary phase consists of Aquapore Butyl 7 micron. The mobile phases are water (with trifluoroacetic acid) and acetonitrile (with trifluoroacetic acid). The injections are 20 µl of a solution of about 1 mg/cm³ in a 0.1 cm² loop valve. The

flow rate for the analyses is adjusted between 1 cm^3/min and 4 $\text{cm}^3/\text{min}.$ The pressure is about 180 bars.

The separation conditions are summarized below:

Solvent A Solvent B

Demineralized water 2500 cm³ Acetonitrile for HPLC 2500 cm³ Trifluoroacetic acid 2 cm³ Trifluoroacetic acid 2.5 cm³

Gradient:

Gradient:					
Time in minutes	% of solvent	% of solvent	Flow rate in		
	A	В	cm³/min		
0	60	40	1		
3	60	40	1		
20	0	100	1		
35	0	100	1		
35.1	60	40	4		
36.1	60	40	4		
36.2	60	40	2		
44	60	40	2		

EXAMPLES

A\ SYNTHESES OF THE COMPOUNDS ACCORDING TO THE INVENTION

Example 1: Synthesis of compound (1)

10 dioctadecylcarbamoylmethyl-2-{3-[4-(2-iminotetrahydropyrimidin-1-yl)butylamino]propylamino}acetamide) from the cationic lipid having the condensed formula NH₂(CH₂)₃NH(CH₂)₄NH(CH₂)₃NHCH₂COGlyN[(CH₂)₁₇CH₃]₂ called "lipid B" in the text which follows (whose preparation 15 has been described in patent application WO 98/18185 and whose structure is represented in Figure 1). 0.784 mmol of lipid B is dissolved in 25 cm³ of methanol in a round-bottomed flask equipped with a magnetic bar, and 10.21 mmol of triethylamine are added. A solution of O-methylisourea and sulphuric acid (1.173 mmol) in 5 water (9 cm³) is then slowly poured (5 minutes) over the

mixture. The mixture is kept at 50°C in an oil bath for twenty hours.

Next, the mixture is concentrated to dryness in a rotary evaporator. The dry extract is solubilized with a solution of water (4 cm³), and trifluoroacetic acid (1 cm³). This solution is injected in two portions in preparative HPLC.

The fractions of interest (determined by analytical HPLC) are grouped together, frozen and freeze-dried.

15 194 mg (0.163 mmol) of salified product are thus obtained.

Yield: 20.8%

HPLCanalytical: Rt = 15.99 minutes.

s respectively, 1H each: the 2 NH of the ring); 8.61

(t, J = 5.5 Hz, 1H: NHCO); 8.70 and 9.02 (2 unres. comp., 1H each: the 2 NH).

 $MH^{*} = 846$

Example 2: Synthesis of compound (2) (N-ditetradecylcarbamoylmethyl-2-{3-[4-(2-iminotetrahydropyrimidin-1yl)butylamino]propylamino}acetamide) from the compound
having the condensed formula

NH2(CH2)3NH(CH2)4NH(CH2)3NHCH2COGlyN[(CH2)13]2 called
"lipid C" in the text which follows (whose preparation
has been described in patent application WO 97/18185
and whose structure is represented in Figure 1).

1.036 mmol of lipid C are dissolved in 30 cm² of
methanol in a round-bottomed flask equipped with a
magnetic bar, and 13.13 mmol of triethylamine are

added. A solution of O-methylisourea and sulphuric acid
(1.554 mmol) in water (9 cm³) is then slowly poured
(5 minutes) over the mixture. The mixture is kept at

20 The dry extract is solubilized with a solution of water $(3~\text{cm}^3)$, ethanol $(2~\text{cm}^3)$ and trifluoroacetic acid $(0.5~\text{cm}^3)$. This solution is injected in preparative HPLC.

50°C in an oil bath for about twenty hours. The mixture is then concentrated to dryness in a rotary evaporator.

The fractions of interest (determined by analytical 25 HPLC) are grouped together, frozen and freeze-dried. 218 mg (0.2022 mmol) of salified product are finally obtained.

Yield: Y = 19.5%

 $MH^{+} = 734$

HPLCanalytical: Rt = 10.76 minutes.

 ^{1}H NMR spectrum (400 MHz, (CD3) $_{2}SO$ d6, δ in ppm): 0.88 (t, J = 7 Hz, 6H: CH₃ of the fatty chains); from 1.15 to 1.40 (mt, 44H: central (CH2)11 of the fatty chains); 1.45 and from 1.50 to 1.65 (2 mts, 2H each: 1 CH2 of each fatty chain); 1.59 (mt, 4H: the 2 central CH_2 of the butyl); 1.91 and 1.97 (2 mts, 2H each: central CH2 of the propyls); from 2.85 to 3.10 (mt, 10H: the 2 NCH2 of the butyl - the 2 NCH_2 of one of the 2 propyls - and 1 of the 2 NHC, of the other propyl); 3.23 and from 3.30 to 3.50 (2 mts, 5H and 1H respectively: the other NCH2 of the other propyl and NCH2 of the fatty chains); 3.79 (unres. comp., 2H: NCH₂CON); 4.03 (d, J = 5 Hz, 2H:CONCHOCON of the glycyl); 7.27 and from 8.40 to 9.30 15 (broad s and unres. comp. respectively, 2H and 4H: $\mathrm{NH_2}^+$ CF3COO; NH+ CF3COO and =NH); 7.88 and 8.61 (s and broad s respectively, 1H each: NHC=N and CONH respectively).

Example 3: Synthesis of compound (3) (2-(3-{4-[3-(4,5-diphydro-1H-imidazol-2-ylamino)propylamino}-N-ditetradecylcarbamoylmethylacetamide) from lipid C (see
Example 2 and Figure 1 for its structure).

0.36 mmol of 2-methylmercapto-2-imidazolinium iodide is dissolved in 0.36 cm³ of 1 N sodium hydroxide in a

25 round-bottomed flask equipped with a bubbler and a magnetic bar. 0.36 mmol of lipid C, previously dissolved in 1.44 cm³ of 1 N sodium hydroxide, 5 cm³ of water and 4 cm³ of ethanol, is added to this solution.

The mixture is kept stirring until the evolution of methyl mercaptan stops (24 hours). The mixture is then concentrated to dryness in a rotary evaporator. The dry extract is solubilized with a solution of water (4 cm 3),

5 ethanol (4 cm 3) and trifluoroacetic acid (0.5 cm 3). This solution is injected in two portions in preparative HPLC.

The fractions of interest (determined by analytical HPLC) are grouped together, frozen and freeze-dried.

10 213 mg (0.1727 mmol) of salified product are finally obtained.

Yield: Y = 48%

HPLCanalytical: Rt = 8.90 minutes.

 1H NMR spectrum (400 MHz, (CD₃)₂SO d6 with addition of a few drops of CD₃COOD d4, δ in ppm): 0.87 (t, J = 7 Hz, 6H: CH₃ of the fatty chains); from 1;15 to 1.40 (mt, 44H: central (CH₂)₁₁ of the fatty chains); 1.45 and 1.55 (2 mts, 2H each: 1 CH₂ of each fatty chain); 1.65 (mt, 4H: the 2 central CH₂ of the butyl); from 1.80 to 1.95

20 (mt, 4H: central CH₂ of the propyls); from 2.80 to 3.05
(mt, 10H: the 2 NCH₂ of the butyl - the 2 NCH₂ of one of
the 2 propyls - and 1 of the 2 HCH₂ of the other
propyl); 3.24 (mt, 6H: the other NCH₂ of the other
propyl and NCH₂ of the fatty chains); 3.56 (s, 2H:

25 NCH_2CON); 3.62 (s, 4H: $NCH_2CH_2CH_2N$); 4.02 (d, J = 5 Hz, 2H: $CONCH_2CON$ of the glycyl).

 $MH^{+} = 777$

Example 4: Synthesis of compound (4) (2-(3-{bis[3-(4,5-dihydro-1H-imidazol-2-ylamino)propyl]amino}propyl-amino)-N-ditetradecylcarbamoylmethylacetamide) by the method for synthesizing "building blocks".

I) SYNTHESIS OF BOC-GLY-DITETRADECYLAMINE (a) The group Gly, whose amines are protected with Boc groups (10 mmol), and the ditetradecylamine (10 mmol) are introduced into a 250 ml round-bottomed flask, and 100 cm3 of dichloromethane are added. The mixture is 10 stirred until complete dissolution is obtained. 30 mmol of N-ethyldiisopropylamine (DIEA) and 11 mmol of benzotriazol-1-yloxytrisdimethylamine phosphonium (BOP) are then added. The pH is kept at 10 by means of the DIEA, and the mixture is stirred for 2 hours. When the 15 reaction is complete, (monitored by CLC and/or HPLC), the dichloromethane is evaporated off and the solid obtained is taken up in ethyl acetate (300 cm3). The organic phase is washed with a solution of potassium sulphate (4 times 100 cm3), of sodium carbonate (4 times 20 100 cm^3), and of sodium chloride (4 times 100 cm^3). The organic phase is then dried over magnesium sulphate, filtered and evaporated under vacuum. The product (a) is obtained with a yield of 93%.

 $\underline{\text{TLC}}$: $R_f = 0.9 \text{ (CHCl}_3/\text{MeOH, 9:1)}$

25 MH+ = 567

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II) SYNTHESIS OF [Z-NH(CH2)3]2-N-(CH2)3-NH-Boc-CH2-COOH (b) 1) Synthesis of NC-(CH2)2-NH-Boc-CH2-COOH (c) The amine of N-(cyanoethyl)glycine (0.1 mol/amine, commercial) is solubilized in 1 N sodium hydroxide (200 cm3/amine) and dioxane (200 cm3). The solution is stirred on an ice bath and then a solution of O-(t-butoxucarbonyl)2 or of pchlorobenzyloxycarbonyl (ClZ, 0.14 mol/amine) in 200 cm3 of dioxane is added dropwise. The pH is kept at a value greater than 9. The mixture is then stirred at about 20°C overnight. The dioxane is evaporated under vacuum and then the mixture is acidified to pH 3 with the aid of a potassium sulphate solution. The insoluble matter is extracted with ethyl acetate (3 times 100 cm3) and then washed with a sodium chloride solution (2 times 100 cm3). The organic phase is dried over magnesium sulphate, filtered and evaporated under vacuum. The product (c) of formula NC-(CH2)2-NH-Boc-CH2COOH is obtained with a yield of 98%. 2.0 TLC: $R_f = 0.66$ (CHCl₃/MeOH, 8:2) $MH^{+} = 229$ 2) Synthesis of NH2-(CH2)3-NH-Boc-CH2-COOH (d)

50 mmol of product (c) of formula NC-(CH2)2-NH-Boc-CH2-COOH are introduced into a 1 litre stainless steel autoclave. A solution of 10 cm3 of ethanol (95%) and of 3.3 q of sodium hydroxide (80 mol) is prepared at the same time in a beaker. When the

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sodium hydroxide has dissolved, 2 cm3 of raney Nickel on carbon are introduced. The autoclave is closed. The initial hydrogenation pressure is about 52 bar, and it decreases to about 48.5 bar overnight at room temperature (20°C). The suspension is filtered on paper, the filter is washed with ethanol (4 times 25 cm3), and the filtrates are concentrated to dryness under vacuum. The product (d) is obtained which is used without further purification in the next stage. TLC: $R_f = 0.12$ (CHCl₃/MeOH, 6:4)

 $MH^{+} = 233$

3) Synthesis of $[NC(CH_2)_2]_2-N-(CH_2)_3-NH-Boc-CH_2-COOH(e)$ The product (d) of formula $NH_2-(CH_2)_3-NH-Boc-CH_2-$ COOH (0.05 mol) and sodium hydroxide (0.1 mol) are solubilized in 150 cm3 of water, in a roundbottomed flask. The solution is cooled on an ice bath. Acrylonitrile (0.12 mol) is slowly poured in, with vigorous stirring, while the temperature of the mass is kept below 20°C. The reaction mixture is kept overnight at room temperature (20°C). The mixture is then kept at 50°C for 2 hours. The solvent is evaporated under vacuum and then the mixture is acidified to pH 3 with a solution of potassium sulphate. The insoluble matter is extracted with ethyl acetate (3 times 200 cm3), and then washed with a sodium chloride solution (2 times 100 cm3). The organic phase is

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dried over magnesium sulphate and then filtered and evaporated under vacuum. The "crude material" is optionally purified on a silica column. The product (e) is obtained with a yield of 50%.

5 $\underline{\text{TLC}}$: $R_f = 0.75$ (CHCl₃/MeOH, 6:4) $MH^+ = 339$

4) Synthesis of [Z-NH(CH₂)₃]₂-N-(CH₂)₃-NH-Boc-CH₂-COOH (b) The product (e) of formula $[NC(CH_2)_2]_2-N-(CH_2)_3-NH-$ Boc-CH2-COOH (50 mmol) is introduced into a 1 litre stainless steel autoclave. A solution of 10 cm3 of ethanol (95%) and of 3.3 g of sodium hydroxide (80 mol) is prepared at the same time in a beaker. When the sodium hydroxide has dissolved, this solution is introduced into the autoclave. A nitrogen stream is passed through the autoclave and 2 cm3 of Raney Nickel on carbon are introduced. The autoclave is closed. The initial hydrogenation pressure is about 52 bar, and it decreases to about 48.5 bar overnight at room temperature (20°C). The suspension is filtered on paper, the filter is washed with ethanol (4 times 25 cm3), and the filtrates are concentrated to dryness under vacuum. A white solid is obtained which is used without further purification after TLC analysis. TLC: $R_f = 0.14$ (CHCl₃/MeOH, 6:4)

25 <u>TLC</u>: $R_t = 0.14$ (CHCl₃/MeOH, 6:4)

The solid obtained above is solubilized in 1 N sodium hydroxide (200 cm³/amine) and dioxane (200 cm³). The solution is stirred on an ice bath

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and then a solution of (t-butoxycarbonyl)₂O or of p-chlorobenzyloxycarbonyl (0.14 mol/amine) in 200 cm³ of dioxane is then added dropwise. The pH is kept at a value greater than 9. The mixture is then stirred at room temperature (20°C) overnight. The dioxane is evaporated under vacuum and then the mixture is acidified to pH 3 with the aid of a potassium sulphate solution. The insoluble matter is extracted with ethyl acetate (3 times 100 cm³) and then washed with a sodium chloride solution (2 times 100 cm³). The organic phase is dried over magnesium sulphate, filtered and evaporated under vacuum. The products are analysed by TLC and/or HPLC.

The crude product is purified on a silica column (dichloromethane/methanol, 8:2).

The product (b) is obtained with a yield of 66% relative to the product (d).

 $\overline{\text{LLC}}$: $R_f = 0.42$ (CHCl₃/MeOH, 6:4)

 $\underline{MH}^* = 615$

III) SYNTHESIS OF [Z-NH(CH₂)₃]₂-N-(CH₂)₃-NH-Boc-CH₂- $\label{eq:coglyn} \mbox{[(CH₂)}_{13}-CH_{3}\mbox{]}_{2} \mbox{ (f)}$

Product (a) whose amines are protected with Boc groups (1 mmol) is introduced into a round-bottomed flask

25 equipped with a magnetic bar. 30 cm³ of trifluoroacetic acid at 4°C are added and then the solution is stirred for one hour. When the reaction is complete (monitored by TLC and/or HPLC), the trifluoroacetic acid is

evaporated under vacuum and then the product is dried by coevaporation with 3 times 30 cm³ of ethyl ether. <u>HPLC</u>: $R_t = 12.86$ min, $(H_2O/MeCN: 3 min [40/60], 3-20 min [0/100], 35 min [0/100].$

- 5 The produced obtained (Gly-ditetradecylamine, 10 mmol) and the product (b) (10 mmol) are introduced into a 250 cm³ round-bottomed flask, dichloromethane (100 cm³) is added and the mixture is stirred until complete dissolution is obtained. 30 mmol of N-ethyldiisopropyl-
- 10 amine (DIEA) and 11 mmol of BOP hexafluorophosphate are then added. The pH is kept at 10 by means of DIEA and the mixture is stirred for two hours. When the reaction is complete (monitored by TLC and/or HPLC), the dichloromethane is evaporated and the solid obtained is
- 15 taken up in ethyl acetate (300 cm³). The organic phase is washed with a potassium sulphate solution (4 times 100 cm^3), of sodium carbonate (4 times 100 cm^3), and of sodium chloride (4 times 100 cm^3). The organic phase is dried over magnesium sulphate, filtered and evaporated
- 20 under vacuum. The products are used without further purification. The product (f) is obtained with a yield of 75% after purification on a silica column (dichloromethane/methanol, 8:2).
 - TLC: $R_f = 0.86 \text{ (CHCl}_3/\text{MeOH, 8:2)}$
- 25 HPLC: Rt = 17.44 min, (H2O/MeCN: 3 min [40/60], 3-20 min
 [0/100], 35 min [0/100]).

IV) SYNTHESIS OF $[NH_2(CH_2)_3]_2-N-(CH_2)_3-NH-Boc-CH_2-COGlyN[(CH_2)_{13}-CH_3]_2$ (g)

Product (f), whose amines are protected, is introduced into a round-bottomed flask equipped with a magnetic

- 5 bar and dissolved in 10 cm³ of methanol per gram of product. Palladium on carbon (10%, 1 g/g of product) and ammonium formate (1 g/g of product) are added at room temperature. Hydrogenolysis is monitored by HPLC. After two hours, the reaction is complete, the mixture
- 10 is filtered and the filter washed with three times
 10 cm² of methanol per gram of product. Double-distilled
 water is added and the solution is frozen and freezedried, or the filtrate is concentrated to dryness and
 the solid is taken up in ethyl acetate (300 cm²). The
- organic phase is washed with a sodium carbonate solution (twice 100 cm³), and a sodium chloride solution (twice 100 cm³), and then it is dried over magnesium sulphate, filtered and evaporated under vacuum. The products are analysed by HPLC and are used without
- 20 further purification. The product (g) is obtained with a yield of 40% relative to the product (f). <u>HPLC</u>: $R_t = 9.62 \text{ min}$, (H_2O/MeCN : 3 min [40/60], 3-20 min [0/100], 35 min [0/100]).

MH+:: 795

25 V) SYNTHESIS OF COMPOUND (4)

Product (g), which contains the primary amine to be modified (1 mmol/amine) is solubilized in dichloromethane (10 cm³) and then 2-methylthio-

imidazoline hydriodide (1.2 mmol/amine) and triethylamine (3 mmol/amine) are added. The mixture is stirred at room temperature (20°C) until the evolution of methyl sulphide stops. At the end of the reaction (monitored by HPLC), the dichloromethane is evaporated

(monitored by HPLC), the dichloromethane is evaporated under vacuum.

The product obtained, whose amines are protected by Boc groups (1 mmol) is introduced into a round-bottomed flask equipped with a magnetic bar. 30 ${\rm cm}^3$ of

- 10 trifluoroacetic acid at 4°C are added and then the solution is stirred for one hour. When the reaction is complete (monitored by TLC and/or HPLC), the trifluoroacetic acid is evaporated under vacuum and then the product is dried by coevaporation with 3 times
 15 30 cm³ of ethyl ether.
 - The product obtained is purified by preparative HPLC and the fractions analysed by HPLC. The compound (4) according to the present invention is thus obtained with a yield of 34%.
- 20 <u>HPLC</u>: $R_t = 10.07$ min, ($H_2O/MeCN$: 3 min [40/60], 3-20 min [0/100], 35 min [0/100]).
 - 1 H NMR spectrum (400 MHz, (CD₃)₂SO d₆ at a temperature of 383 K, d in ppm): 0.92 (t, J = 7 Hz, 6H: CH₃ of the fatty chains); from 1.25 to 1.45 (mt, 44H: (central
- 25 (CH₂)₁₁ of the fatty chains); 1.57 (mt, 4H : 1 CH₂ of
 each fatty chain); from 1.70 to 1.90 (mt, 6H : central
 CH₂ of the propyls); from 2.50 to 3.40 (mt, 16H : the
 2 NHC₂ of the propyls and the NCH₂ of the fatty chains);

3.68 (s, 8H : the 2 NCH_2CH_2N); 3.72 (broad s, 2H : NCH_2CON); 4.06 (s, 2H : $CONCH_2CON$ of the glycyl). MH^{+} : 831

Example 5: Synthesis of compound (5):

- 5 (N-Ditetradecylcarbamoylmethyl-2-{3-[3-(1,4,5,6-tetrahydropyrimidin-2-ylamino)propylamino]propylamino}-acetamide) by the method of synthesis of "building blocks".
 - I) SYNTHESIS OF BOC-GLY-DITETRADECYLAMINE (a)
- 10 The procedure is carried out in the same manner as in the preceding example. Product (a) is obtained with a yield of 93%.

TLC: $R_f = 0.9$ (CHCl₃/MeOH, 9:1)

MH+: 567

- 15 II) SYNTHESIS OF Z-NH(CH2)3-N-Boc-(CH2)3-N-Boc-CH2-COOH (b)
 - 1) Synthesis of NC-(CH₂)₂-NH-Boc-CH₂-COOH (c)
 The procedure is carried out in the same manner as above in Example 4. Product (c) is obtained with a yield of 98%.
- 20 TLC: Rf = 0.66 (CHCl₃/MeOH, 8:2)

MH+: 229

2) Synthesis of NH₂-(CH₂)₃-NH-Boc-CH₂-COOH (d)

Product (d) is obtained in the same manner as above in Example 4.

25 $\underline{\text{TLC:}}$ R_f = 0.12 (CHCl₃/MeOH, 6:4) MH*: 233

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3) Synthesis of NC(CH2)2-N-Boc-(CH2)3-NH-Boc-CH2-СООН (e)

Product (d) (0.05 mol) and sodium hydroxide (0.1 mol) are solubilized in 150 cm3 of water in a round-bottomed flask. The solution is cooled on an ice bath. Acrylonitrile (0.05 mol) is slowly poured in, with vigorous stirring, while the temperature of the mass is kept below 20°C. The reaction mixture is kept overnight at room temperature (20°C).

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The solvent is evaporated under vacuum and then the mixture is acidified to pH 3 with a potassium sulphate solution. The insoluble matter is extracted with ethyl acetate (3 times 200 cm3), and then washed with a sodium chloride solution (twice 100 cm3). The organic phase is dried over magnesium sulphate and then filtered and evaporated under vacuum. The product obtained is optionally purified on a silica column.

2.0 The product obtained (0.1 mol/amine) is solubilized in 1 N sodium hydroxide (200 cm3/amine) and dioxane (200 cm3). The solution is stirred on

an ice bath and then a solution of (Boc),0 or of p-chlorobenzyloxycarbonyl (0.14 mol/amine) in

200 cm3 of dioxane is added dropwise. The pH is kept at a value greater than 9. The mixture is then stirred at room temperature (20°C) overnight.

The dioxane is evaporated under vacuum and then

the mixture is acidified to pH 3 with the aid of a potassium sulphate solution. The insoluble matter is extracted with ethyl acetate (3 times 100 cm³) and then washed with a sodium chloride solution (twice 100 cm³). The organic phase is dried over magnesium sulphate, filtered and evaporated under vacuum. The products are analysed by TLC and/or HPLC.

Product (e) is thus obtained with a yield of 93%. $\overline{\text{LLC}}\colon \ R_f \ = \ 0.75 \ (\text{CHCl}_3/\text{MeOH}, \ 8:2)$

MH+: 386

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4) Synthesis of Z-NH-(CH₂)₃-N-Boc-(CH₂)₃-N-Boc-CH₂COOH (b)

Product (e) (50 mmol) is introduced into a onelitre stainless steel autoclave. A solution of
10 cm³ of ethanol (95%) and of 3.3 g of sodium
hydroxide (80 mol) is prepared at the same time in
a beaker. When the sodium hydroxide has dissolved,
this solution is introduced into the autoclave. A
nitrogen stream is passed through the autoclave
and 2 cm³ of Raney Nickel on carbon are introduced.
The autoclave is closed. The initial hydrogenation
pressure is about 52 bar and it decreases to about
48.5 bar overnight at room temperature (20°C). The
suspension is filtered on paper, the filter is
washed with ethanol (4 times 25 cm³) and the
filtrates are concentrated to dryness under

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vacuum. A white solid is obtained which is used without further purification after TLC analysis.

TLC: $R_f = 0.14$ (CHCl₃/MeOH, 6:4)

The product obtained (0.1 mol/amine) is solubilized in 1 N sodium hydroxide (200 cm3/amine) and dioxane (200 cm3). The solution is stirred on an ice bath and then a p-chlorobenzyloxycarbonyl solution (0.14 mol/amine) in 200 cm3 of dioxane is added dropwise. The pH is kept at a value greater than 9. The mixture is then stirred at room temperature (20°C) overnight. The dioxane is evaporated under vacuum and then the mixture is acidified to pH 3 with the aid of a potassium sulphate solution. The insoluble matter is extracted with ethyl acetate (3 times 100 cm3) and then washed with a sodium chloride solution (twice 100 cm3). The organic phase is dried over magnesium sulphate, filtered and evaporated under vacuum. The product obtained is purified on a silica

column (dichloromethane/methanol, 9:1). The products are analysed by TLC and/or HPLC. Product (b) is obtained with a yield of 32% relative to product (d).

TLC: $R_f = 0.63$ (CHCl₃/MeOH, 9:1)

25 MH+: 523

- III) SYNTHESIS OF 2-methylsulphanyl-1,4,5,6-tetrahydropyridmidine (f)
- 3,4,5,6-Tetrahydro-2-pyrimidinethiol (0.0103 mol) is loaded into a round-bottomed flask, with stirring and
- 5 under a nitrogen stream, and 5 cm³ of methanol and 0.65 cm³ of methyl iodide (0.0105 mol) are added. The mixture is heated under reflux for 1 hour and is then allowed to cool to room temperature (20°C). The product is precipitated by addition of 5 cm³ of ethyl ether. The
- 10 precipitate is filtered and then washed with ethyl ether. The product is then dried overnight at a pressure of 34 mbar.
 - 1.5 g (0.0041 mol) of product (VI) are obtained, that is to say a yield of 40%.
- 15 TLC: $R_f = 0.25$ (CHCl₃/MeOH, 9:1)

MH+: 131

- d) SYNTHESIS OF Z-NH(CH $_2$) $_3$ -N-Boc(CH $_2$) $_3$ -N-Boc-CH $_2$ -COGlyN[CH $_2$) $_{13}$ -CH $_3$] $_2$
- Product (a), whose amines are protected with Boc groups

(q)

- 20 (1 mmol) is introduced into a round-bottomed flask equipped with a magnetic bar. 30 cm³ of trifluoroacetic acid at 4°C are added and then the solution is stirred for one hour. When the reaction is complete (monitored by TLC and/or HPLC), the trifluoroacetic acid is
- 25 evaporated under vacuum and then the product obtained is dried by coevaporation with 3 times 30 cm³ of ethyl ether.

<u>HPLC:</u> $R_t = 12.86 \text{ min}$, $(H_2O/MecN: 3 \text{ min } [40/60], 3-20 \text{ min } [0/100], 35 \text{ min } [0/100])$.

The product obtained (10 mmol) and the product (b) (10 mmol) are introduced into a 250 cm³ round-bottomed

- 5 flask. Dichloromethane (100 cm³) is added and the mixture is stirred until complete dissolution is obtained. 30 mmol of DIEA and 11 mmol of BOP are then added. The pH is kept at 10 by means of the DIEA and the mixture is stirred for two hours. When the reaction
- 10 is complete (monitored by TLC and/or HPLC), the dichloromethane is evaporated and the solid obtained is taken up in ethyl acetate (300 cm³). The organic phase is washed with a solution of potassium sulphate (4 times 100 cm³), of sodium carbonate (4 times
- 15 100 cm³), and of sodium chloride (4 times 100 cm³). The organic phase is dried over magnesium sulphate, filtered and evaporated under vacuum. The products are used without further purification.

After purification on a silica column

20 (dichloromethane/methanol, 8:2), product (g) is obtained with a yield of 85%).

TLC: $R_f = 0.9$ (CHCl₃/MeOH, 9:1)

<u>HPLC</u>: $R_t = 19.79$ in, $(H_2O/MeCN)$: 3 min [40/60], 3-20 min [0/100], 35 min [0/100]).

25 V) SYNTHESIS OF NH₂(CH₂)₃]₂-N-Boc-(CH₂)₃-NH-Boc-CH₂COGlyN[(CH₂)₁₃-CH₃]₂ (h)
Product (g) is introduced into a round-bottomed flask
equipped with a magnetic bar and dissolved in 10 cm³ of

methanol/q of product. Palladium on carbon (10%, 1 q/q of product) and ammonium formate (1 g/g of product) are added at room temperature (20°C). The hydrogenolysis is monitored by HPLC. After two hours, the reaction is 5 complete, the mixture is filtered and the filter is washed with 3 times 10 cm3 of methanol/g of product. Double-distilled water is added and the solution is frozen and freeze-dried, or the filtrate is concentrated to dryness and the solid is taken up in 10 ethyl acetate (300 cm3). The organic phase is washed with a solution of sodium carbonate (twice 100 cm3), and a solution of sodium chloride (twice 100 cm3) and then it is dried over magnesium sulphate, filtered and evaporated under vacuum. The products are analysed by 15 HPLC and are used without further purification. Product (h) is obtained with a yield of 93% relative to product

TLC: $R_f = 0.42$ (CHCl₃/MeOH, 6:4)

<u>HPLC</u>: $R_t = 14.66 \text{ min}$, $(H_2O/MeCN: 3 \text{ min } [40/60], 3-20 \text{ min}$

20 [0/100], 35 min [0/100]).

MH+: 838

(q).

VI) SYNTHESIS OF COMPOUND (5)

Product (h) containing the primary amine to be modified (1 mmol/amine) is solubilized in dichloromethane

25 (10 cm³) and then product (f) (1.2 mmol/amine) and triethylamine (1.3 mmol/amine) are added. The mixture is stirred at room temperature (20°C) until the evolution of methyl sulphide stops. At the end of the

reaction (monitored by HPLC), the dichloromethane is evaporated under vacuum.

The product obtained is purified by preparative HPLC and the fractions analysed by HPLC. Compound (5) is thus obtained with a yield of 38%.

<u>HPLC</u>: $R_t = 8.42 \text{ min}$, $(H_2O/MeCN: 3 \text{ min } [40/60]$, 3-20 min [0/100], 35 min [0/100]).

 $\frac{^{1}\text{H NMR spectrum}}{\text{J} = 7 \text{ Hz}}$ (400 MHz, (CD₃)₂SO d₆ δ in ppm): 0.86 (t,

- 10 1.35 (mt, 44H : central (CH₂)₁₁ of the fatty chains);
 1.44 and 1.53 (2 mts, 2H each : 1 CH₂ of each fatty
 chain); from 1.80 to 2.00 (mt, 6H : central CH₂ of the
 propyls and CH₂ of 1,4,5,6-tetrahydropyrimidine); from
 2.80 to 3.10 (mt, 10H : NCH₂ of the propyls and NCH₂ of
- 15 1,4,5,6-tetrahydropyrimidine); from 3.14 to 3.45 (mt :
 the 6H corresponding to the =NCH₂ of 1,4,5,6-tetrahydro pyrimidine and to the =NCH₂ of the fatty chains); 3.81
 (unres. comp., 2H : NCH₂CON); 4.04 (d, J = 5 Hz, 2H:
 CONCH₂CON of the glycyl); 7.89 = 8.62 8.75 and 9.01
- 20 (4 unres. comp., 8H in total : the exchangeables and OH of the CF,COOH).

MH+: 720

Example 6: Synthesis of compound (6): N-Dioctadecylcarbamoylmethyl-2-{3-[3-(1,4,5,6-tetrahydropyrimidin-2ylamino)propylamino]propylamino}acetamide) by the
method of synthesis of "building blocks".

I) SYNTHESIS OF BOC-GLY-DITETRADECYLAMINE (a)

The procedure is carried out in the same manner as in the preceding example. Product (a) is obtained with a yield of 93%.

5 TLC: $R_f = 0.9$ (CHCl₃/MeOH, 9:1)

MH+: 567

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- II) SYNTHESIS OF Z-NH(CH2)3-N-Boc-(CH2)3-N-Boc-CH2-COOH (b)
 - 1) Synthesis of NC-(CH2)2-NH-Boc-CH2-COOH

The procedure is carried out in the same manner as above in Example 5. Product (c) is obtained with a vield of 98%.

TLC: Rf = 0.66 (CHCl₃/MeOH, 8:2)

MH+: 229

- 2) Synthesis of NH2-(CH2)3-NH-Boc-CH2-COOH (d)
- 15 Product (d) is obtained in the same manner as above in Example 5.

TLC: $R_f = 0.12$ (CHCl₃/MeOH, 6:4)

MH+: 233

3) Synthesis of NC(CH2)2-N-Boc-(CH2)3-NH-Boc-CH2-

20 соон (e)

> The procedure is carried out in the same manner as above in Example 5. Product (e) is thus obtained with a yield of 93%.

TLC: $R_f = 0.75$ (CHCl₃/MeOH, 8:2)

MH+: 386

25

10

4) Synthesis of Z-NH- $(CH_2)_3$ -N-Boc- $(CH_2)_3$ -N-Boc- CH_2 COOH (b)

The procedure is carried out in the same manner as above in Example 5. A white solid is obtained which is used without further purification after a TLC analysis.

TLC: $R_f = 0.14$ (CHCl₃/MeOH, 6:4)

The product obtained is used in the same manner as above so as to protect the terminal amine with a benzyloxycarbonyl group. Product (b) is thus obtained with a yield of 32% relative to product (d).

 $\underline{\text{TLC:}} \ R_{\text{f}} = \text{0.63 (CHCl}_{3}/\text{MeOH, 9:1})$ $MH^{+}: \ 523$

15 III) SYNTHESIS OF 2-methylsulphanyl-1,4,5,6-tetrahydropyridmidine (f)

The procedure is carried out in the same manner as above in Example 5. 1.5 g (0.0041 mol) of product (f) are thus obtained, that is to say a yield of 40%.

20 TLC: $R_f = 0.25$ (CHCl₃/MeOH, 9:1)

MH+: 131

IV) SYNTHESIS OF Z-NH(CH₂)₃-N-Boc(CH₂)₃-N-Boc-CH₂- COGlyN[(CH₂)₁₇-CH₃]₂ (g)

The procedure is carried out in the same manner as

25 above in Example 5. Product (a) is thus obtained whose Boc groups have been cleaved.

<u>HPLC</u>: $R_t = 19.44$ min, ($H_2O/MeCN$: 3 min [40/60], 3-20 min [0/100], 35 min [0/100]).

This product obtained is used in the same manner with product (b) as above in Example 5. After purification on a silica column (dichloromethane/methanol, 8:2), product (g) is obtained with a yield of 84%.

5 TLC: $R_f = 0.9$ (CHCl₃/MeOH, 9:1)

<u>HPLC</u>: $R_t = 23.95 \text{ min}$, ($H_2O/MeCN$: 3 min [40/60], 3-20 min [0/100], 35 min [0/100]).

V) SYNTHESIS OF NH₂ (CH₂) $_3$] $_2$ -N-Boc-(CH₂) $_3$ -NH-Boc-CH $_2$ -COGlyN[(CH $_2$) $_1$ 7-CH $_3$] $_2$ (h)

10 The procedure is carried out in the same manner as above with Example 5. Product (h) is obtained with a yields of 73% relative to product (g).

TLC: $R_f = 0.28$ (CHCl₃/MeOH, 6:4)

 $\underline{\text{HPLC}}\colon$ R_{t} = 20.59 min, (H₂O/MeCN: 3 min [40/60], 3-20 min

15 [0/100], 35 min [0/100]).

MH+: 838

VI) SYNTHESIS OF COMPOUND (6)

The procedure is carried out in the same manner as above in Example 5. Compound (6) is thus obtained with

20 a yield of 68%.

<u>HPLC</u>: $R_t = 15.83$ min, ($H_2O/MeCN$: 3 min [40/60], 3-20 min [0/100], 35 min [0/100]).

 1 H NMR spectrum (500 MHz, (CD₃)₂SO d₆ δ in ppm): 0.88 (t, J = 7 Hz, 6H : CH₃ of the fatty chains); from 1.15 to

25 1.35 (mt, 60H: central $(CH_2)_{15}$ of the fatty chains); 1.46 and 1.54 (2 mts, 2H each: 1 CH_2 of each fatty chain); from 1.80 to 2.00 (mt, 6H: central CH_2 of the propyls and CH_2 of 1,4,5,6-tetrahydropyrimidine); from 2.85 to 3.05 (mt, 10H: NCH₂ of the propyls and NCH₂ of 1,4,5,6-tetrahydropyrimidine); from 3.15 to 3.45 (mt: the 6H corresponding to the =NCH₂ of 1,4,5,6-tetrahydropyrimidine and to the NCH₂ of the fatty chains); 3.81 (unres. comp., 2H: NCH₂CON); 4.04 (d, J = 5 Hz, 2H = CONCH₂CON of the glycyl); 7.88 = 8.61 - 8.74 and 8.99 (4 unres. comp., 8H in total: the exchangeables and OH of the CF₃COOH).

MH+: 832

10 B\USE OF THE TRANSFECTION AGENTS ACCORDING TO THE INVENTION

Example 7: preparation of nucleolipid complexes

This example illustrates the preparation of nucleolipid complexes according to the invention.

The compound used in this example is compound (1) in solution in chloroform. 10 nmol quantities of compound (1) (that is to say 11.8 μg) per μg of DNA were used. In some cases, a neutral co-lipid, Cholesterol or DOPE, is previously mixed with the 20 compound. A fine lipid film forms when the chloroform is evaporated with the aid of a slight stream of argon, and then it is rehydrated in a mixture of 5% dextrose and 10 mM sodium chloride overnight at 4°C. The samples are then treated with ultrasound for 5 minutes, heated 25 at 65°C for 30 minutes and finally treated again with ultrasound for 5 minutes. Lipid suspensions are thus obtained which are stored at 4°C until they are used.

The DNA used is the plasmid pXL2774

(Figure 2) in solution in a mixture of 5% dextrose and 20 mM sodium chloride at a concentration of 0.5 mg/ml or 1.0 mg/ml. The plasmid pXL2774 has the following 5 characteristics:

- level of endotoxins less than 50 EU/mg,
- level of supercoiled DNA greater than 60%,
- content of RNA, that is to say of mRNA, tRNA and ribosomal RNA (determined by HPLC) less than 5%,
 - level of chromosomal DNA less than 1%,
 - protein content less than 1%,
 - osmolarity less than 15 mosmol/kg.

The nucleolipid complexes according to the invention are prepared by rapidly mixing equal volumes of DNA solution and lipid suspension as described above. The quantity of compound complexed with the DNA varies from 0.5 nmol/µg of DNA to 12 nmol/µg of DNA.

Example 8: behaviour of the complexes formed at different charge ratios

This example illustrates the behaviour of the nucleolipid complexes according to the invention at different charge ratios. The impact of the addition of a neutral co-lipid is also illustrated.

The size of the complexes was first of all

25 analysed by measuring the hydrodynamic diameter by
dynamic light scattering (Dynamic Laser Light

Scattering) with the aid of a Coulter N4plus apparatus.

The samples are diluted 20-fold in a solution

containing 5% dextrose and 20 mM sodium chloride in order to avoid multiple diffusions. The effect of the cycloamidine group, of the lipid composition and of the charge ratio on the size of the nucleolipid complexes 5 according to the invention was thus studied.

Three possible phases can be distinguished
when the charge ratio between compound (1) according to
the invention and the DNA is increased. These three
phases determine the therapeutic potential of
10 compound (1). Figure 3 illustrates these 3 phases for
compound (1). The same behaviour can be observed for

other compounds according to the invention.

At a low charge ratio, the DNA is not saturated with compound (1). Naked DNA still remains,

and the complexes are negatively charged overall. The particles are small in size (between 100 and 300 nm). This phase is called "Phase A".

The fact that the DNA is not completely saturated with compound (1) means that the DNA is not completely

protected by it. The DNA can therefore be subjected to degradation by enzymes (DNAses). Moreover, since the complexes are negative overall, the crossing of the cell membrane is difficult. For these reasons, the nucleolipid complexes of phase A are of a much lower efficiency in transfection.

At an intermediate charge ratio, the DNA is completely saturated with compound (1), and the complexes are neutral or slightly positive overall. This phase is unstable because the ionic repulsions are minimal and a "crosslinking" phenomenon may occur. The size of the particles is well above the limit of detection by dynamic light scattering (much greater than 3 \mummam). This unstable phase is called "phase B". Such a size of complexes is not suited to uses by injection. However, this does not necessarily mean that the complexes are inactive in phase B, but they are only in a formulation which is not appropriate for their injection for pharmaceutical purposes.

At a relatively high charge ratio, the DNA is oversaturated with compound (1), and the complexes are positive overall. Because of the strong repulsions between the positive charges, this phase is stable. It is designated by the name "phase C". Unlike phase A, the nucleolipid complexes are in a form such that the DNA is very well protected against enzymes, and their overall positive charge facilitates the crossing of the cell membrane of anionic nature. The phase C complexes are therefore particularly suited to use for the transfer of nucleic acids into cells.

In addition to the cycloamidine group of the compound according to the invention, the use of a neutral co-lipid has a strong impact on the stability

25 of the complexes, as is illustrated in Figure 3. The co-lipids added are either DOPE (cationic lipid/DOPE = 3/2), or cholesterol (cationic lipid/cholesterol = 3/1). In general, the addition of the neutral co-lipid

increases the instability of the complexes, which causes an increase in the quantity of compound required to obtain phase C. This is very clearly illustrated in Figure 3 when the charge ratio at which phase C is obtained in the presence and in the absence of co-lipid is compared.

It should be noted that the values of the charge ratio which delimit the three phases A, B and C depend on the compound used. Thus, these values can 10 vary very widely from one compound to another.

Finally, the affinity of the compound for the DNA as a function of the charge ratio was studied. For that, the reduction in fluorescence after the addition of 3 μg of ethidium bromide (EtBr) was measured.

15 Indeed, the replacement of the ethidium bromide of the DNA by the compound is an indication of binding to the DNA.

The formulation used is diluted 20-fold to a final concentration of 25 μ g of DNA/ml. The relative 20 fluorescence measured for naked DNA is defined as being 100%. The level of binding with compound (1) is represented by the reduction in the relative fluorescence of the sample. Figure 3 shows that the fluorescence decreases when the charge ratio increases, which means that a greater quantity of compound (1) is available to bind to the DNA (the more the fluorescence decreases, the more a large quantity of compound binds to the DNA until saturation is reached).

In this manner, it has been shown that the affinity of compound (1) according to the invention for the DNA is determined by the cycloamidine group, but not by the addition of a co-lipid.

5 Example 9: transfection in vitro with compound (1)

This example illustrates the capacity of compound (1) according to the invention to transfect DNA into cells in vitro, compared with nonformulated DNA.

24-well microplates are inoculated with
60,000 HeLa cells (ATCC) per well, and transfected
24 hours later. Complexes containing 1 μg of DNA are
diluted in 0.5 ml of DMEM culture medium (Gibco/BRL) in
the absence of serum, and then added to each well. The
15 cells are incubated at 37°C for 4 hours. The medium
containing the complexes is then removed and replaced
with a mixture of DMEM and 10% foetal calf serum. Next,
the cells are again cultured for 24 hours. Finally, the
cells are lysed and tested using a luciferase test kit
20 (Promega) and a Dynex MLX luminometer.

The results indicated in Figure 4 underline
the difference between the performance of the naked DNA
compared with the compound (1)/DNA complexes of the
invention which are completely saturated: no luciferase
activity could be detected (sensitivity of the
apparatus less than 1 pg per well) after transfection
in vitro of naked DNA, whereas the gene transfer

activity of the complexes according to the invention varies from 200 pg/well to 8000 pg/well.

This example therefore clearly shows the advantageous use of compound (1) according to the 5 invention for the transfection of cells in vitro.

Example 10: transfection in vitro with compounds (3), (5) and (6)

This example illustrates the capacity of compounds (3), (5) and (6) according to the invention to transfect DNA into cells in vitro, compared with nonformulated DNA.

The transfection is carried out according to the preceding protocol of Example 9, into HeLa cells.

The results are illustrated in Figures 5, 6 and 7. It is thus observed that these 3 compounds have a good transfection level in vitro.

Example 11: transfection in vivo of compound (1)

This example illustrates the capacity of compound (1) according to the invention to transfect

20 DNA into cells in vivo, compared with nonformulated DNA and with lipid A having the condensed formula

NH2 (CH2) 3NH (CH2) 4NH (CH2) 3NHCH2COArgN [CH2) 17CH3] 2 described in Application WO 97/18185 and whose structure is represented represented in Figure 1.

25 The gene transfer in vivo was performed on Balb/C mice by intramuscular and intravenous administration. The formulations which were compared are formulations of naked DNA, formulations containing

lipid A, or formulations containing compound (1) according to the invention.

In the case of intramuscular injections, each mouse received 30 µl of formulation containing 15 µg of DNA

5 in the anterior muscle of the tibia. The tissues are recovered 7 days after the injection, they are frozen and stored at -80°C while waiting to perform the luciferase activity tests. The measurements of luciferase activity are carried out as in Example 8.

10 In the case of injections by the intravenous route, each mouse received 200 µl of formulation containing 50 µg of DNA. The tissues are recovered in this case 24 hours after the injection and then frozen and stored

The results of gene transfer in vivo are presented in Figure 8 and Figure 9. The ratio between compound (1) and the DNA is 10 nmol/ μ g of DNA. The ratio between lipid A and the DNA is 4 nmol/ μ g of DNA.

in the same manner as above.

Figure 8 illustrates the *in vivo* activity in

20 the muscle of compound (1) according to the invention
compared with naked DNA and with lipid A. It is
observed that the levels of luciferase activity are
equivalent between naked DNA and compound (1), the
latter having, in addition, a highly improved activity

25 compared with lipid A. The transfer mechanisms involved
appear to be different between naked DNA and the use of
compound (1) according to the present invention.
Indeed, the complexes according to the invention used

do not contain free DNA (phase C) and furthermore, their results in vitro are considerably greater than those for naked DNA.

Figure 9 compares the activity of

5 compound (1) according to the invention, of naked DNA
and of lipid A, by the intravenous route and by the
intramuscular route.

It is observed that the transfection efficiency is roughly equivalent by the intravenous 10 route for lipid A and for compound (1). On the other hand, by the intramuscular route, the transfection efficiency of compound (1) according to the invention is quite considerably greater than that of limid A.

Compared with naked DNA, compound (1)

15 exhibits transfection by the intravenous route, in addition to transfection by the intramuscular route which is at least equivalent.

It therefore appears that the nucleic acid transfer efficiency in vivo with compound (1) according 20 to the invention is greater overall than that with lipid A which is a known cationic lipid and that of nonformulated DNA.

Finally, it appears that the complexes according to the invention have the advantage, compared 25 with transfection of naked DNA, of protecting the DNA from degradation by nucleases, thus contributing to a significant improvement in the stability of the formulations. The compounds of the present invention

can also be used to protect DNA from damage during freeze-drying, improving here again the stability of the formulations.

Example 12: transfection in vivo of compounds (5) and 5 (6)

This example illustrates the capacity of compounds (5) and (6) to transfect nucleic acid *in vivo* in an efficient manner.

The same protocol as in the preceding example

10 is used. Figure 10 shows that compound (5) and

compound (6), formulated in a 0.25:1 charge ratio with

DNA without co-lipid, exhibit a transfection level

in vivo greater than or equal to maked DNA 48 hours

after i.m. injection.

The following table gives the results obtained with compounds (5) and (6) in various formulations:

Compound	Compound/	Co-lipid	RLU/	pg/	Route of
	DNA		lung	lung	adminis-
	charge				tration
	ratio				
Compound (5)	6/1	DOPE (1:1)	254.9	1611.3	i.v.
Compound (5)	8/1	DOPE (1:1)	535.2	3558.1	i.v.
Compound (5)	0.5/1	Chol. (3:1)	209.6	1330.5	i.m.
Compound (5)	0.5/1	DOPE (1:1)	155.8	974.6	i.m.
Compound (6)	6/1	-	175.1	1098.7	i.v.
Compound (6)	5/1	DOPE (1:1)	407.7	2700.8	i.v.
Compound (6)	0.5/1	DOPE (1:1)	1768.7	13005.4	i.m.

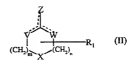
CLAIMS

 Compounds, in D, L or DL form, and salts thereof, of general formula (I):

$$CA - Rep - R$$
 (I)

5 for which:

 $\ensuremath{\Phi}$ CA represents a cycloamidine group and its mesomeric forms of general formula (II):



10 for which:

- ullet m and n are integers, independent of each other, of between 0 and 3 inclusive and such that m+n is greater than or equal to 1,
- \bullet R_{1} represents a group of general formula (III):

$$- \left[(CH_2)_p - Y \right]_q (*)$$
 (III)

15

for which p and q are integers, independent of each other, of between 0 and 10 inclusive, Y represents a carbonyl, amino, methylamino or methylene group, it being possible for Y to have different meanings within the different groups [(CH₂)_p-Y], and (*) represents either a hydrogen atom or is the site for bonding to the group Rep,

it being understood that R_1 may be bonded to any atom of general formula (II), including Z, and that there is a single group R_1 in formula (II),

- X represents a group NR₂ or CHR₂, R₂ being either a
 hydrogen atom or the bonding to the group R₁ as defined above,
 - The group v represents:

*1st case: a group of general formula(IV):

10 for which W' represents CHR \square or NR \square , and R" and R \square represent, independently of each other, a hydrogen atom, a methyl, or the bonding to the group R₁ as defined above, or

*2nd case: a group of general formula (V):

15

for which W' represents CHR \square or NR \square , and R' and R \square represent, independently of each other, a hydrogen atom, a methyl or the bonding to the group R₁ as defined above,

$$- \left\{ \begin{array}{c} N - (CH)_r \end{array} \right\}_{t}^{0} \qquad (VI)$$

whose nitrogen atom is attached to the atoms X, V, W or Z or to the substituent Y of the group R_1 depending on the cases, and

- 5 t is an integer between 0 and 8 inclusive,
 - r is an integer between 0 and 10 inclusive, it being possible for r to have different meanings within the different groups $-NR_4-(CH)_{\pi^-}$,
- R_3 , which may have different meanings within the different groups NR_4 -(CH) $_2R_3$, represents a hydrogen atom, a methyl group or a group of general formula (VII):

$$\frac{-\left\{(CH_2)_s - N\right\}_{U}H}{R_s} \qquad (VII)$$

for which u is an integer between 1 and 10 inclusive, s

15 is an integer between 2 and 8 inclusive which may have
different meanings within the different groups
-(CH₂)_s-NR₅, and R₅ is a hydrogen atom, a group CA as
defined above, it being understood that the groups CA
are independent from each other and may be different,

20 or a group of general formula (VII), it being
understood that the groups of general formula (VII) are

independent of each other and may have different meanings,

- R₄ is defined in the same manner as R₃ or represents a group CA as defined above, it being understood that the
 groups CA are independent of each other and may be different, and
- * either a group of formula NR₆R₇ for which R₆ and R₇ represent, independently of each other, a hydrogen atom or an optionally fluorinated, linear or branched, saturated or unsaturated aliphatic radical containing 1 to 22 carbon atoms, with at least one of the two
- 15 substituents R_6 or R_7 different from hydrogen and the other containing between 10 and 22 carbon atoms,
 - * or a steroid derivative,
 - * or a group of general formula (VIII):

$$-[NH-(CH2)x]y Q (VIII)$$

20 for which x is an integer between 1 and 8 inclusive, y is an integer between 1 and 10 inclusive, and either Q represents a group C(O)NR₆R₇ for which R₆ and R₇ are as defined above, or Q represents a group C(O)R₈ for which R₈ represents a group of formula (IX):

$$\begin{array}{c|c}
R_6 & R_6 \\
N & N & O \\
N & O & R_9
\end{array}$$
(IX)

for which z is an integer between 2 and 8 inclusive, and R_9 is an optionally fluorinated, saturated or unsaturated aliphatic radical containing 8 to 22 carbon 5 atoms, or a steroid derivative, and the two substituents R_6 are, independently of each other, as defined above,

or R_{8} represents a group -O-R9 for which R_{9} is as defined above.

- 2. Compounds according to claim 1, characterized in that the group R_1 is bonded either to Z or to V, on the one hand, and to the group Rep, on the other hand, via Y.
- Compounds according to claim 1,
 characterized in that the cycloamidine head CA of formula (II) comprises 5, 6, 7 or 8 members.
- 4. Compounds according to claim 1, characterized in that R_3 represents a hydrogen atom or a methyl and R_4 is as defined in claim 1, or R_3 and R_4 20 present in formula (VI) represent hydrogen atoms, or R_4 is a hydrogen atom and R_3 is a group of formula (VII) in which R_5 represents a group CA.

- 5. Compounds according to claim 1, characterized in that in formula (V), p and q are chosen, independently of each other, from 2, 3 or 4.
 - 6. Compounds according to claim 1,
- 5 characterized in that the groups R₆ and R₇ are identical or different and each represent optionally fluorinated, linear or branched, saturated or unsaturated aliphatic chains containing 10 to 22 carbon atoms.
 - 7. Compounds according to claim 1,
- 10 characterized in that the groups R_6 and R_7 are identical or different and each represent optionally fluorinated, linear or branched, saturated or unsaturated aliphatic chains containing 12, 14, 16, 17, 18 or 19 carbon atoms.
- 15 8. Compounds according to claim 1, characterized in that when R is a steroid derivative, the said steroid derivative is chosen from cholesterol, cholestanol, $3-\alpha-5$ -cyclo- $5-\alpha$ -cholestan- $6-\beta$ -ol, cholic acid, cholesteryl formate, chotestanyl formate,
- 3α,5-cyclo-5α-cholestan-6β-yl formate, cholesterylamine, 6-(1,5-dimethylhexyl)-3a,5a-dimethylhexadecahydrocyclopenta[a]cyclopropa[2,3]cyclopenta[1,2-f]naphthalen-10-ylamine or cholestanylamine.
 - 9. Compounds according to claim 1 of
- 25 formulae:

10

Compound (6)

10. Method of preparing the compounds
according to claims 1 to 9, characterized in that the
synthesis of the building blocks carrying the
cycloamidine function(s) is carried out and then these

building blocks are grafted onto lipids equipped with spacers.

- 11. Method of preparing the compounds according to claims 1 to 8, characterized in that the 5 synthesis of the analogous lipopolyamines is carried out and then the cyclization into cycloamidine groups is carried out.
 - 12. Composition, characterized in that it comprises at least one compound of general formula (1).
 - 13. Composition according to claim 12, characterized in that it comprises a compound of general formula (1) and a nucleic acid.
- 14. Composition according to claims 12 or13, characterized in that it comprises, in addition,15 one or more adjuvants.
 - 15. Composition according to claim 14, characterized in that the adjuvant(s) are one or more neutral lipids containing two fatty chains.
 - 16. Composition according to claim 15,
- 20 characterized in that the neutral lipids are natural or synthetic lipids which are zwitterionic or lacking ionic charge under physiological conditions, chosen for example from dioleoylphosphatidylethanolamine (DOPE), oleoylpalmitoylphosphatidylethanolamine (POPE),
- 25 di-stearoyl, -palmitoyl, -mirystoylphosphatidylethanolamines as well as their derivatives which are N-methylated 1 to 3 times, phosphatidylqlycerols,

diacylglycerols, glycosyldiacylglycerols, cerebrosides (such as in particular galactocerebrosides), sphingolipids (such as in particular sphingomyelins) or asialogangliosides (such as in particular asialoGM1 and 5 GM2).

- 17. Composition according to claim 14, characterized in that the adjuvant is a compound involved directly or otherwise in the condensation of the nucleic acid.
- 18. Composition according to claim 17, characterized in that said adjuvant is derived as a whole or in part from a protamine, a histone or a nucleolin and/or from one of their derivatives, or consists, as a whole or in part, of peptide units 15 (KTPKKAKKP) and/or (ATPAKKAA), it being possible for the number of units to vary between 2 and 10, and to be repeated continuously or otherwise.
- 19. Composition according to claims 12 to 18, characterized in that it contains, in addition, one 20 or more nonionic surfactant(s) in a sufficient quantity to stabilize the size of the particles of nucleolipid complexes.
 - Composition according to claims 12 to 19, characterized in that it comprises a vehicle which is pharmaceutically acceptable for an injectable formulation.

- 21. Composition according to claims 12 to 19, characterized in that it comprises a vehicle which is pharmaceutically acceptable for an application to the skin and/or the mucous membranes.
- 22. Composition according to claim 13, characterized in that the said nucleic acid is a deoxyribonucleic acid or a ribonucleic acid.
- 23. Composition according to claim 22, characterized in that the said nucleic acid comprises

 10 an expression cassette consisting of one or more genes of therapeutic interest under the control of one or more promoters and of a transcriptional terminator which are active in the target cells.
- 24. Use of a compound according to one of 15 claims 1 to 9, to manufacture a medicament for treating diseases.
- 25. Use of a compound according to one of claims 1 to 9, to manufacture a medicament for treating diseases by transfer of nucleic acids into cells by the 20 intramuscular route.
 - 26. Method of transferring nucleic acids into cells comprising the following steps:
 - bringing the nucleic acid into contact with a compound of general formula (1) as defined above, to
- 25 form a nucleolipid complex, and
 - (2) bringing the cells into contact with the nucleolipid complex formed in (1).

- 27. Method of transferring nucleic acids into cells according to claim 26, characterized in that the said nucleic acid and/or the said compound are previously mixed with one or more adjuvants.
- 28. Method of treating diseases by administration of a nucleic acid encoding a protein or which can be transcribed into a nucleic acid capable of correcting the said diseases, the said nucleic acid being combined with a compound of general formula (I).

ABSTRACT

5 Novel compounds relating to the lipopolyamine family and comprise at least one cyclic amidine function are provided. The novel compounds form nucleolipid complexes when contacted with nucleic acids. The compounds are useful for transferring nucleic acids into cells by in vitro, ex vivo or in vivo methods.

PCT/FR99/00740

1/10

FIG. 1/10

lipid B

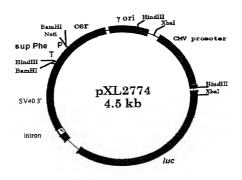
$$\mathsf{H}^\mathsf{M} \sim \mathsf{M} \sim \mathsf{$$

lipid C

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2/10

FIG. 2/10

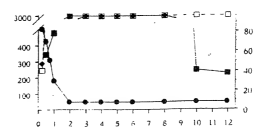


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FIG. 3/10

Compound 1

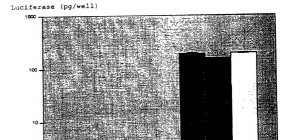


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Compound (1)

4/10

FIG. 4/10



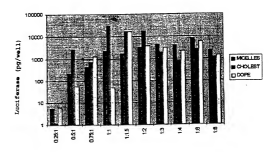
Naked DNA

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5/10

FIG. 5/10

Compound (3), in vitro (HeLa cells)

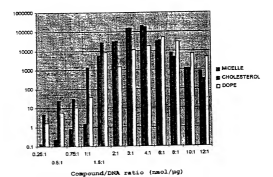


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6/10

FIG. 6/10

Compound (5), in vitro (HeLa cells)

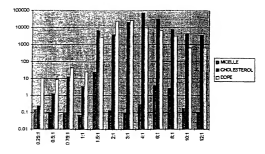


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7/10

FIG. 7/10

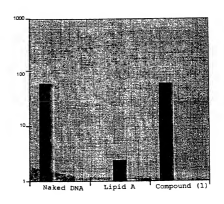
Compound (6), in vitro (heLa cells)



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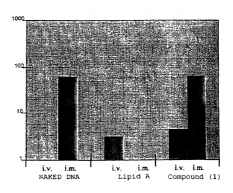
FIG. 8/10



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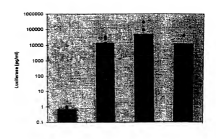
FIG. 9/10



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FIG. 10/10





DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

As a below named inventor(s), I/We hereby declare that:

My/Our residence(s), post office address(es) and citizenship(s) are as stated below my name(s).

I/We verily believe I am/we are the original, first and sole/joint inventor(s) of the subject matter which is claimed and for which a patent is sought on the invention entitled: NEW AGENTS FOR TRANSFERRING NUCLEIC ACIDS, COMPOSITIONS CONTAINING THEM AND THEIR USES and the specification of which is attached hereto (Attorney Docket No.) was filed on 02 Oct • 2000 as U.S. Application Number (check one) and was amended on (if applicable).
was described and claimed in PCT Int'l Application Number filed on and as amended under PCT Article 19 on I/We hereby state that I/We have reviewed and understand the contents of the above identified specification, including the claims as amended by any amendment referred to above. I/We acknowledge the duty to disclose to the U.S. Patent and Trademark Office all information known to me/us to be material to patentability as defined in 37 C.F.R. 1.56. I/We hereby claim foreign priority benefits under Titte 35, United States Code §119(a)-(d) or 365 (b) of any foreign application(s) for patent or inventor's certificate, or 365(a) of any PCT international application which designated teast one other country other than the United States of America, listed below and having a filing date before that of the application on which priority is claimed. I/We have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed: 02 April 1998 / Day/Month/Year Filed FR98/04121 Number France

Foreign Appln(s):

PCT/FR99/00740

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Foreign

Priority:

Prior

П

I/We hereby claim the benefit under Title 35, United States Code §119(e) of any United States Provisional application(s) listed below:

30 March 1999 / Day/Month/Year Filed

Number Filing Date

Country

I/We hereby claim the benefit under Title 35, United States Code §120 or 365(c) of any United States application(s) or international application designating the United States listed below and, insofar as the subject matter of each of the international application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35. United States application in the manner provided by the first paragraph of Title 35. United States application in the manner provided by the first paragraph of Title 35. United States Code §112. [We acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations §1.56 which became available between the filing date of the prior application and the national or PCT international filing date of this application.

Application Serial No. Filing Date Status (Patented, Pending)

I/We hereby appoint the attorneys and/or agents associated with the Customer No.(s) provided below as my/our attorneys and/or agents with full power to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith:

> Customer No.: 005487 Send Correspondence to:

> > Karen I. Krupen

Aventis Pharmaceuticals Products Inc. Rt. 202-206/Patent Dept.

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I/We hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Title 18, United States code §1001, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.



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